Choosing from the conversation smorgasbord of a cocktail party

Barbara Shinn-Cunningham

At a rollicking party, conversations ebb and flow all around. Although the noise level may make it difficult to hear some exchanges, there are often multiple discussions that could be understood. This is very convenient if you find yourself in dull conversation about, say, motor homes, while the group behind you is sharing some racy gossip about one of your colleagues. In such situations, you can satisfy social expectations by monitoring what is happening in front of you, even though you are really processing the chatter from behind. This real-life example illustrates the various factors that can affect the ability to communicate in social settings. Early psychoacoustic work focused on understanding how a competing masker sound interferes with the early sensory representation of information about an important message, and how much of that message must be represented cleanly for it to be understood. While such details determine whether or not you understand a message when there is only one source demanding your attention, such as in a phone conversation, that is not how you operate when you navigate a busy street or argue some controversial issue with impassioned colleagues at a faculty meeting. How your brain focuses on different conversations depending on your interests governs what knowledge you take away from a cocktail party as surely as the does the fidelity of the sensory representation.

A View of the Precedence Effect through the Owl's Ears

Terry Takahashi, Brian Nelson, Caitlin Baxter

In nature, sounds of interest are followed by reflections from nearby surfaces. Yet, for short delays, the sound arriving directly from the source dominates perception. This phenomenon, “localization dominance” (LD), is part of the precedence effect. To better understand the neural mechanisms of the precedence effect, we conducted neurophysiological and psychoacoustical experiments in the barn owl (Tyto alba), with a direct sound (ds) that overlaps with a simulated echo.

We found that LD requires deep amplitude modulations (AMs) when the onset and offset delays are removed. Without these AMs, the owl localized the leading and lagging sounds with equal frequency, in spite of the fact that the lagging carrier was a delayed copy of the lead.

To explain these observations, we developed the “envelope model”. When there are two sound sources with overlapping amplitude spectra, the frequency-specific binaural cues are biased toward values corresponding to the momentarily-louder source. Neurons in the owl's auditory space map discharge if, as the binaural cues begin to approximate those of its spatial receptive field, the amplitude is rising. The envelope model computes the relative incidences of this conjunction to predict the number of spikes evoked by the lead and lag sources. The model suggests that LD occurs because a peak in the lead obscures the rising edge of the corresponding peak in the lag.

Reflected sounds and independent, competing sounds are on a continuum of similarity to the ds. When the lead and lag sounds have different envelopes (but identical, but delayed carriers) there are no corresponding peaks. In such case, the strengths of the lead and lag sounds' neural representations, which can be computed by the envelope model, are determined by the arbitrary shapes of the envelopes. Results show that the owl localizes the sound predicted to have the stronger representation.

Our results suggest that 1.) The temporal relationship between the lead/lag pair is determined not from the carrier, but from the ongoing envelope disparities. 2.) LD is due to the superposition of the waveforms in the ears and the selectivity of neurons for upward AMs, suggesting echoes need not be actively suppressed. 3.) Lead/lag pairs and a pair of independent sounds are represented on the space map according to the same principles.
Neural Bases of Concurrent Sound Segregation in Primate Auditory Cortex
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The ability to perceptually separate concurrent sounds (e.g., voices) is critical for successful hearing in complex acoustic environments (e.g., a cocktail party or noisy cafeteria). Neural mechanisms underlying concurrent-sound segregation are poorly understood and are thought to involve central auditory processes. Here, we present results of electrophysiological studies in primary auditory cortex (A1) of awake monkeys that aim to shed light on these neural mechanisms. We examine neural correlates of two perceptual phenomena reflecting key principles of concurrent sound segregation: (1) "pop-out" of mistuned components in harmonic complex sounds, exemplifying segregation based on inharmonicity, and (2) segregation of concurrent harmonic complex sounds (e.g., vowels) based on a difference in their fundamental frequency (F0).

In relation to (1), we find that responses to harmonic complex tones (HCTs) containing a mistuned component are enhanced in neural populations that are tuned to the frequency of the mistuned component, relative to responses evoked by 'in-tune' HCTs. This response enhancement parallels the increased perceptual salience of mistuned harmonics reported in human psychoacoustic studies. Mistuned sounds also evoke temporal discharges which are phase-locked to lower frequency "beats" in the stimuli. Thus, a local increase in firing rate or a difference in temporal response pattern in A1 may contribute to concurrent sound segregation based on inharmonicity.

In relation to (2), using a stimulus design employed in auditory-nerve studies (Larsen et al., 2008), we found that neural responses in A1 convey sufficient spectral ('rate-place') and temporal ('phase-locking') information reflecting the harmonic structure and F0s of two simultaneous and spectrally overlapping HCTs to enable their perceptual segregation. Lower harmonics (1-6) of the HCTs were better resolved in neural 'rate-place' representations than higher harmonics (7-12), consistent with psychoacoustic findings in humans. Furthermore, introducing an onset asynchrony between the HCTs enhanced the neural representation of their harmonics, consistent with the improved ability to perceptually segregate asynchronous sounds. F0s below 350 Hz were also represented in temporal discharges phase-locked to the periodicities of the HCTs. Findings suggest that A1 contains sufficient spectral and temporal information for extracting the pitch of HCTs via harmonic templates and periodicity detectors, and for segregating concurrent HCTs based on a difference in F0. Improved understanding of central mechanisms contributing to concurrent sound segregation may facilitate the development of more effective approaches toward alleviating deficits in speech and music perception in hearing-impaired listeners.
Cortical Encoding of Auditory Objects at the Cocktail Party
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A visual scene is perceived in terms of its constituent visual objects. Similar ideas have been proposed for the analogous case of auditory scene analysis, though their hypothesized neural underpinnings have not yet been established. Here, we investigate how auditory objects are individually represented in auditory cortex, using magnetoencephalography (MEG) to record the neural responses of human listeners. Subjects selectively listen to one of two competing streams, in a variety of auditory scenes. The auditory streams may overlap spatially, spectrally, or both.

First we demonstrate that attentional gain does not act globally on the entire auditory scene, but rather acts differentially on the separate auditory streams. This stream-based attentional gain is then used as a tool to individually analyze the different neural representations of the competing auditory streams.

In the acoustically richest example, subjects selectively listen to one of two competing speakers mixed in a single channel. Individual neural representations of the speech of each speaker are observed in auditory cortex, with each being selectively phase locked to the rhythm of the corresponding speech stream, and from which can be exclusively reconstructed the temporal envelope of that speech stream. The neural representation of the attended speech, originating in posterior auditory cortex, dominates the responses. Critically, when the intensities of the attended and background speakers are separately varied over a wide intensity range, the neural representation of the attended speech adapts only to the intensity of that speaker, but not to the intensity of the background speaker. This demonstrates object-level intensity gain control in addition to the object-level attentional gain.

Overall, these results indicate that concurrent auditory objects, even if spectrally overlapping and not resolvable at the auditory periphery, are indeed neurally encoded individually as separate objects, in auditory cortex.

Cochlear implants in real-life auditory scenes: limitations and opportunities
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Cochlear implants are up to now the most successful man-made interfaces to the neural system. The auditory nerve is stimulated electrically which leads to a partial restoration of hearing and auditory perception for persons with severe hearing impairment. Speech understanding of cochlear implant (CI) recipients in quiet environments can be very good, but considerably worse in more real-life and adverse listening situations.

Although modern CIs use up to 22 stimulation channels, the information transfer is still very limited for the perception of fine spectro-temporal details to allow the perception of music and speech communication in common real-life auditory scenes. The limitations have become clear after years of research. Major limitations are related to reduced spectral resolution and the lack of matching of the coding of temporal and spectral aspects of pitch.

However, temporal aspects of the input sound, such as the speech envelope and periodicity can very well be transmitted in CIs. Examples will be given of enhanced representation of these features in the signal coding leading to improved directional hearing and speech perception in challenging listening scenes. Furthermore, some signal processing approaches introduce speech enhancements in noisy conditions at the cost of significant signal distortions. These distortions are evaluated as detrimental for sound quality in normal hearing and hearing impaired, but are hardly noticeable by most cochlear implant recipients. This opportunity for further improvements in auditory perception in CI will be discussed.

Dynamic faces speed up vocal processing in the auditory cortex
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The dynamics and deformations of the face during speech lead to a variety of visual motion cues related to the auditory components of speech; they are integral to face-to-face communication. In noisy, real world environments, visual cues provide considerable intelligibility benefits to the perception of auditory speech and faster reaction times. We are currently investigating the neural correlates of this behavioral advantage by investigating what role the auditory cortex of macaque monkeys plays in the audiovisual integration of their "coo" vocalizations. Monkeys detected visual, auditory or audiovisual
coo vocalizations produced by monkey avatars in a background of noise (~60 dB) as fast and as accurately as possible. The loudness of the auditory vocalization (i.e. signal to noise ratio or SNR) relative to the noise background was varied. Decreasing SNR of the auditory-only vocalizations led to an increase in reaction times (RTs). Audiovisual RTs were faster than RTs to both auditory- and visual-only conditions. Under these behavioral conditions, we recorded spiking activity and local field potential (LFPs) in the lateral belt auditory cortex. We found that, at least in part, the activity in auditory cortex co-varied with behavior. First, a decrease in SNR of the auditory-only vocalization increased latency and decreased the magnitude of spiking responses. Second, surprisingly, we found that spiking responses were faster for audiovisual compared to auditory vocalizations—a parallel with decreasing reaction times. Spiking responses were however absent to visual-only vocalizations — suggesting sub-threshold effects of visual input. Thus, we analyzed the LFP responses to identify the network dynamics that could mediate this speed-up. We found that the inter-trial phase coherence in the 10 – 30 Hz band of the LFP was suppressed for audiovisual compared to auditory vocalizations. Suppression was maximal for the largest SNR and decreased with the decrease in SNR. This suggests that visual input into auditory cortex changes the state of network dynamics in this frequency band. We also found that the onset of visual mouth motion led to an increase in the inter-trial phase coherence in the 10 – 30 Hz band immediately after visual onset. These results suggest that during the detection of visual vocalizations, visual cues speed up and alter the dynamics of the circuits in auditory cortex that process vocalizations.

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A Temporal Integration Mechanism Enhances Frequency Selectivity Between Brainstem Inputs and Inferior Colliculus Neurons
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Mammals can accurately resolve frequency components in sounds, a task that is essential for sound recognition. Despite its importance, there is little direct evidence for how frequency selectivity is preserved or newly created across auditory structures. Here we demonstrate a dramatic enhancement in frequency selectivity between putative input (PIN) and target inferior colliculus neurons (ICN) that is created through temporal integration of the converging inputs. PIN receptive fields are broadly tuned and exhibit temporally delayed and spectrally interleaved excitation and inhibition not present in IC. A radical sharpening of tuning is accomplished by temporal integration of the converging inputs. A neuron model accurately replicates the finding and demonstrates that this integration degrades timing precision but enhances spectral selectivity through interference of spectrally in- and out-phase IC inputs. This contrasts current models that require local inhibition to enhance selectivity and supports a fast computational strategy to enhance frequency selectivity in IC.

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Making a case for integration time-constants: Do sparse representations scale hierarchically?
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Neural codes that are minimally redundant and sparse have been proposed as efficient strategies for representing natural sounds. Prevailing theories hold that neural responses to sound shift hierarchically from highly redundant dense codes in the auditory periphery to a selective sparse codes in cortex. On the other hand, computational models that use sparse coding principles can account for receptive field features in auditory nerve (Lewicki 2002), midbrain (Carlson et al. 2012), and cortex (Klein et al 2003). These seemingly conflicting views can be reconciled if one considers the neural integration times for phase-locked activity, which increase systematically from the auditory nerve to auditory cortex. Here we demonstrate that most inferior colliculus (ICC) neurons in the cat respond with a single precise action potential within their spectrotemporal response field (STRF), which defines their integration time. Population activity was likewise sparse with few neurons active for a typical integration time and neuron-to-neuron correlations where minimal (5% of pairs) and strictly confined to 1/3 octave. Responses from primary auditory cortex (A1) are similarly sparse if each structure is analyzed at their respective integration time scales (10 msec ICC; 50 msec A1). This indicates that sparse representations scale hierarchically with neural integration times and that measures of sparseness should be referenced on the time-scales relevant to the sound feature represented at each stage of the auditory pathway.
Associative Learning Enhances Population Coding in the Avian Auditory Cortex by Inverting Inter-Neuronal Correlation Patterns

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Cortical encoding of acoustic signals involves the coordinated activity of millions of neurons. Although learning can alter encoding in individual cortical neurons, it is unknown how this plasticity extends to neural populations. Correlated inter-neuronal activity, a property not observable in single neuron responses, can greatly impact population encoding and is thus a potential target for learning driven plasticity. To test this possibility, we measured the simultaneous activity of multiple auditory cortical neurons in songbirds trained to recognize natural segments of song. We show that learning inverts the canonical positive relationship between correlated signal (tuning function similarity) and correlated noise between pairs of neurons. This inversion enhances the ability of neural populations to discriminate between those song segments that are relevant for behavior. These results reveal the inter-neuronal correlation relationship as a novel target for learning-dependent plasticity.

Sensitivity and Invariance to Temporal Transformations of Vocalizations in the Primary Auditory Cortex

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One of the central tasks of the mammalian auditory system is to represent information about acoustic communicative signals, such as vocalizations. However, the neuronal mechanisms underlying vocalization encoding, and sensitivity to its temporal transformations, in the central auditory system are poorly understood. To learn how the auditory cortex encodes information about con-specific vocalizations, we presented a library of natural and transformed ultra-sonic vocalizations (USVs) to awake rats, while recording neural activity in A1 using chronically implanted multi-electrode probes. The vocalizations were presented in their original form, reversed temporally, dilated or compressed. For responses of each neuron to each stimulus group, we fitted a simple, yet novel predictive model: a generalized linear-non-linear model (GLNM) that takes the frequency modulation and single-tone amplitude as the only two input parameters. Many neurons reliably and selectively responded to original USVs. Our GLNM accurately predicted their responses to previously unheard USVs. Neurons with high model prediction accuracy were typically tuned to tone pips in the ultra-sonic vocalization frequency range. However, accurate model performance is also reported for neurons that were tuned to lower frequencies. We next tested the changes in neuronal responses to temporally transformed and reversed versions of the original vocalizations. The model prediction accuracy was lower for temporally transformed, rather than original or reversed vocalizations, whether the model was fitted on the original or the transformed vocalizations, whereas the mean firing rate was not affected by the transformations of the stimulus. Our results demonstrate that A1 processes original USVs differentially than transformed USVs, indicating specialization for temporal statistics of the original vocalizations.

Auditory Coding of Individual Vocalizations in Scenes

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Background
Processing communication vocalizations is a crucial, natural function of the auditory system. Central auditory circuits generate representations of vocalizations that lead to perception and guide behavior. An important part of understanding how auditory processing leads to perception of communication vocalizations in complex scenes is explaining mechanisms whereby neural representations of vocalizations are extracted from neural representations of scenes.

Methods
We trained songbirds to discriminate among individual vocalizations and tested their abilities to recognize those vocalizations in auditory scenes at varying signal to noise ratios. We then recorded the responses of single auditory midbrain, primary forebrain and higher forebrain neurons to the same vocalizations at varying intensities and in scenes at varying signal to noise ratios. Using analysis of recorded spike trains, pharmacological and stimulus manipulations, and simulations, we generated and tested a simple neural circuit model that explains the transformation in neural representations of vocalizations and scenes along the ascending auditory pathway.

Results
We found that the neural representation of vocalizations transformed from a dense and redundant code in the midbrain and primary forebrain into a sparse and distributed code in the higher forebrain. Sparse coding neurons were better than dense coding neurons at extracting individual vocalizations from auditory scenes, and generated neural representations of target vocalizations in scenes at signal to noise ratios that matched behavioral detection. A simple neural circuit of fast
excitation and delayed inhibition transforms dense input into sparse output, and facilitates the neural extraction of an individual vocalization from a complex auditory scene.

Conclusions
Our findings suggest that the neural representation of vocalizations transforms significantly between the primary and higher auditory forebrain, from a dense and non-selective code to a sparse, highly selective and distributed code. The higher forebrain sparse code is sensitive to acoustic context, relies on inhibition, and serves as a mechanism for the neural extraction of one vocalization from a complex scene.

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Mechanisms of attention in auditory scene analysis
Mounya Elhilali

The mechanisms by which a complex auditory scene is parsed into coherent objects depend on poorly-understood interactions between task-driven and stimulus-driven attentional processes. It is widely believed that detecting a target amidst ambient noises or competing sounds is not only mediated by top-down processes that direct listeners' attention to the signal of interest; but is also modulated by the listening environment, statistics of the auditory scene and prior information that listeners have or gather about the scene. Here, we explore the role of exogenous (stimulus-driven) processes in forming priors about our listening environments; and their interaction with endogenous (listener-directed) attentional processes. Our findings indicate that listeners operate in an optimal way by integrating all information from both task instructions and stimulus and scene cues in an effective manner. We propose a computational framework that integrates the role of both endogenous and exogenous processes in foreground/background organization and auditory object formation.

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Natural Sound Statistics and Auditory Scene Analysis
Josh McDermott, Eero Simoncelli

Auditory scene analysis involves estimating individual sound sources from the signal that enters the ear, which often contains mixtures of multiple concurrent sources. The segregation of sounds from a mixture is a classic ill-posed problem in perception, and can be solved only with the aid of prior assumptions about the sound sources that occur in the world. In this talk we will present a theoretical framework by which these assumptions about sound sources can be related to natural sound statistics. We view sound segregation as the task of distinguishing actual sound sources from the many possible incorrect alternatives – sound signals that are physically consistent with the mixture received by the ears but which did not occur in the world. Our approach stems from the observation that incorrect estimates of individual sources must tend to be partial mixtures of the true sources. Sound properties that have different values for individual sources compared to mixtures should thus aid segregation, as they could serve to distinguish correct and incorrect source estimates. By comparing the properties of individual sound sources with those of their mixtures, we can assess the theoretical utility of different potential segregation cues. We will discuss the results of such statistical analyses of various corpora of natural sounds, their implications for theories of segregation, and how these implications might be tested experimentally.
A mutation in *PNPT1*, encoding the mitochondrial RNA-import protein PNPase, causes hereditary hearing loss

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**Background**

A subset of nuclear encoded RNAs has to be imported into mitochondria to ensure proper replication and transcription of the mitochondrial genome and hence mitochondrial function. Polynucleotide phosphorylase (PNPase, encoded by *PNPT1*) is one of the few components known to be involved in this poorly characterized process in mammals. In our present study we investigated a consanguineous Moroccan family with autosomal recessive non-syndromic hearing impairment (DFNB70) and identified a homozygous missense mutation in *PNPT1* as the underlying cause of disease.

**Methods**

Family members were evaluated by detailed medical history interviews, a physical examination, and underwent an otological examination and pure-tone audiometry. To identify the chromosomal locus underlying the sensory disorder, we performed genome-wide homozygosity mapping. Coding exons and adjacent splice sites of all positional candidate genes within the linked interval were analysed by PCR amplification of genomic DNA followed by direct Sanger sequencing. RT-PCR was applied to investigate *PNPT1* expression in various murine tissues including the cochlea. In zebrafish the expression of the *PNPT1* ortholog was analyzed by in situ hybridization. To investigate the cellular localization of PNPase within the cochlea, we performed immunohistochemistry on cochleae cross-sections and in organ of Corti whole-mount preparations. The ability of the PNPase mutant to assemble into functional trimers was analyzed by blue-native gel electrophoresis and mitochondrial RNA import assays were conducted in yeast and mammalian cells.

**Results**

By positional cloning we identified a homozygous *PNPT1* missense mutation (c.1424A>G predicting the protein substitution p.Glu475Gly) of a highly conserved PNPase-residue within the second RNase PH-domain in a family with autosomal recessive non-syndromic hearing impairment. The immunohistological staining pattern demonstrated a broad PNPase distribution in the inner ear including the sensory hair cells and spiral ganglion neurons. In vitro analyses in bacteria, yeast, and mammalian cells showed that the identified mutation results in a hypofunctional protein leading to disturbed PNPase trimerization and reduced mitochondrial RNA import.

**Conclusion**

In summary, we elucidated a hereditary disturbance of the mitochondrial RNA import by the identification of a hypofunctional mutation in *PNPT1* in congenital severe hearing impairment, demonstrating the rather unexpected and specific importance of PNPase for auditory function.

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**Gene Expression Profiling of Young and Adult Mouse Cochlea by RNA-Seq in Strains with Normal and Age-Related Hearing Loss**

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**Background**

Age-related hearing loss is the most common sensory deficit that reduces the quality of life in the aged population. Its social impact will become more pronounced as life expectancy increases. Despite the discovery of many deafness genes, pathophysiology of progressive hearing impairment due to aging remains elusive. We hypothesize that gene expression profiling of the cochlea from mouse models with various degrees of age-related hearing loss will reveal the molecular mechanism and inform potential target selection for prevention and treatment.

**Methods**

We performed gene expression profiling of mouse cochlea by next-generation sequencing (RNA-Seq). A multi-factorial design was used. Cochleas from mouse strains with good hearing past one year of age (CBA/CaJ and B6.CAST-
Cdh23<sup>ΔN42</sup>) or with documented age-related hearing loss (Coch<sup>G88E/G88E</sup> and Coch<sup>−/−</sup> in a CBA background and C57BL/6J) were dissected at discrete ages ranging from one week through late adulthood. PolyA selected mRNAs were extracted and the derived cDNA samples were fragmented, indexed, pooled, and sequenced by IlluminaHiSeq. Biological replicates were used for all conditions. Expression levels of all transcripts were analyzed and differential gene expression analyses were performed.

**Results**
We created gene expression profiles of mammalian cochlea at various ages by RNA-Seq. With total reads of at least 30 million, more than 16,000 genes were detectable, and expression levels were highly reproducible. We detect significant systematic differences in gene expression profiles between C57BL/6J and CBA/CaJ strain backgrounds, regardless of age. Comparing mouse models with or without age-related hearing loss of the same genetic background, we find that few genes show statistically significant differential expression at young ages before the onset of hearing impairment, but the number of genes dramatically increases to hundreds at later stages. In addition, hundreds of genes show significant temporal changes on both genetic backgrounds. This postnatal to adult cochlea specific gene expression dataset will eventually be added to the publicly accessible Shared Harvard Inner Ear Database (SHIELD: https://shield.hms.harvard.edu).

**Conclusion**
We have surveyed gene expression in mammalian cochlea by RNA-Seq and identified genes that show age-related differential expression. Systematic differences exist between different genetic backgrounds. Temporal gene expression profiles in the cochlea may suggest candidate targets for prevention and treatment of age-related hearing loss. By providing online access to the fully annotated dataset, we expect it to become a useful resource for the research community.

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**Deep Sequencing Identifies Novel microRNAs in the Mouse Inner Ear**

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**Background**
MicroRNAs (miRNAs) are small non-coding RNAs that regulate gene expression post-transcriptionally. By binding to sequences in the 3’ untranslated region (UTR) of mRNAs, a miRNA can inhibit target mRNAs by translational suppression and mRNA destabilization. miRNAs clearly play an important role in the inner ear, since mutations in miRNAs lead to deafness in humans and mice. Conditional knock-out mouse mutants have shown that miRNAs are vital for development of the sensory neurons, sensory epithelia and inner ear morphogenesis.

**Methods**
To further characterize miRNAs in the mammalian inner ear, we used Illumina deep sequencing to identify novel miRNAs in mouse inner ear sensory epithelia and differentially-expressed miRNAs in cochlea and vestibule. Short RNA molecules were sequenced from cochlear and vestibular sensory epithelia of newborn C57Bl/6 mice by high-throughput DNA sequencing. The reads were aligned to the mature *Mus musculus* miRNA database, allowing one mismatch to occur between a read and the reference, due to isomiRs. The reads that were not mapped to known miRNAs were used to predict novel pre-miRNAs or pri-miRNAs, according to the known characteristic read pattern for each type of novel RNA, using the DARIO algorithm. We selected miRNAs with the following criteria: seed regions conserved between mouse and human, highly expressed in the sensory epithelia, and located in introns of genes expressed in the inner ear due to possible co-expression.

**Results**
A total of 7,732,589 and 8,452,794 small RNAs were found in the cochlear and vestibular samples, respectively. These included miRNAs, snoRNAs, transfer RNAs and ribosomal RNAs. For miRNAs, 440 and 458 were identified in the cochlea and vestibule sensory epithelium, respectively. In addition, we identified three novel mouse miRNAs, qRT-PCR and in situ hybridization validated expression of these miRNAs in the inner ear. Predictions of gene-targtes to these miRNAs were verified by luciferase assays. The interaction between the miRNAs and a number of their targets is being studied.

**Conclusion**
We have generated and analyzed a comprehensive dataset of small RNA sequences from the mouse inner ear. The analysis revealed novel miRNAs with potential regulatory effects on development. Based on the function of the targets,
these miRNA-target pairs may be involved in cell signaling, apoptosis, membrane structure, and conductance of voltage-gated channels, with implications for hearing and mechanisms leading to deafness.

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Genome wide association analysis and expression studies involved in hearing function and age-related hearing loss

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Background

Only few genes have been involved in normal hearing function and age-related hearing loss (ARHL). To reach the goal of increasing our knowledge on the genetic bases of both hearing function and ARHL an integrated strategy has been designed based on: A) Genome Wide Association Studies (GWAS) on different hearing traits (low, medium and high Pure Tone Average, the separated thresholds and the Principal components), B) expression studies in mice using immunohistochemistry and confocal microscopy of genes identified by the GWAS and C) genotype-phenotype relationships

Methods

Three meta-analyses using 3681 samples coming from isolated populations located in Europe (Italy, Croatia and Ukraine), Caucasus (Armenia, Azerbaijan, Georgia) and Central Asia (Uzbekistan, Kazakhstan, Tajikistan and Kirghizstan) were carried out. Two GWAS for hearing quantitative traits (one sex separated), and one for ARHL (qualitative traits) have been performed. In order to confirm our findings, expression studies in wild-type mice (at 4 and 5 days postnatal) using immunohistochemistry and confocal microscopy were then carried out on genes identified by the GWAS. Genotype-phenotype relationships have been performed using mean values for each frequency and comparing the 3 different genotypes of a given SNP

Results

By meta-analysis of data in these populations, some significant and suggestive loci (p<10^-6,10^-7) have been found and some of them map to known hereditary hearing loss loci whose genes still need to be identified (Girotto et al. JMG 2011). A list of 22 strong candidate genes has been defined to be included in the expression studies. 5 of them (Arsg, Slc16a6, Dclk1, Gabrg3, Csmd1) show strikingly specific expression in the cochlea (e.g. at the top of sensory hair cells and in the marginal cells of the stria vascularis) while the other 10 (Ptprd, Grm8, GlyBP, Evi5, Irg1, Rimb2, Ank2, Mbd1, Rbfox1, Cdh13) are located in multiple cell types in the cochlea. As regards the possible genotype-phenotype correlation, an example is reported in Figure 1 where Irg1 and Grm8, being both expressed in many cell types in the cochlea, show similar pattern at all the frequencies in the three genotypes. Moreover, some interesting findings have been obtained comparing phenotype/genotype data with the localization of the expression

Conclusion

Preliminary results prove the useful combination of GWAS, expression studies and genotype/phenotype data to provide new insights into the molecular basis of hearing function and ARHL suggesting new targets for treatment and prevention
**Exome Sequencing Reveals Mutations in ATP6V1B2 as a Cause of Dominant Deafness-Onychodystrophy (DDOD) Syndrome**

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**Background**

Dominant deafness and onychodystrophy syndrome (DDOD syndrome) is characterized by congenital sensorineural hearing loss, dystrophic or absent of nails. Cornical and hypoplastic teeth were also present in some cases. The DDOD syndrome as an autosomal dominant condition was first recognized in 1961 and subsequently identified in five other unrelated families. The genetic etiology of DDOD is not yet determined.

**Methods**

We collected two Chinese DDOD pedigrees. The probands showed identical phenotypes: congenital hearing impairment, absent of all the toe nails, little finger and thumb nails, absent of the distal little finger bones as well as onychodystrophy like malacia and pitting of the middle three finger nails. The whole-exome sequencing was performed in two probands and their parents using the genomic DNA extracted from peripheral blood. The on-target mean depth of exome sequencing was >130 fold, resulting in >99% of the targeted base pairs being sufficiently covered. Single-nucleotide variations and indels of patients and their parents were filtered by dbSNP132, the 1000 Genomes Project, HapMap8 databases and BGI YH database. Candidate genes of inner ear from the Morton Human Fetal cDNA library were also used in the filtering. Immunohistochemistry and specific gene knockdown by using morpholino antisense oligos were carried out to investigate the pathogenic mechanism of DDOD-related gene mutations.

**Results**

Mutations in four genes, ATP6V1B2, ACTG1, PABPC1 and CDC27 were identified in the two affected probands by our analysis strategy. Since the quality scores of PABPC1 and CDC27 SNPs were low, we concentrated on confirmatory sequencing of the variations in the ATP6V1B2, ACTG1 by Sanger sequencing. A common heterozygous de novo mutation 1516 C>T (R506X) in the ATP6V1B2 was verified. While the ACTG1 variant turned out to be a false positive. Conservation analysis of amino acids in nine ATP6V1B2 orthologs indicated the arginine in 506 is highly conserved in evolution. In addition, the 1516 C>T mutation in ATP6V1B2 was not detected in 1053 ethnically matched normal controls. Immunohistochemistry labeling of the ATP6V1B2 in the cochlea identified its expression in the hair cells, spiral ganglia, spiral limbus, interdental cells, spiral prominence and fibrocytes. Mice injected with ATP6V1B2 morpholino antisense oligos into the scala media of cochlea showed significant hearing loss in the injected side, while contralateral control showed normal hearing.

**Conclusion**

Exome sequencing data identified a mutation in the ATP6V1B2 as the cause of DDOD. Preliminary functional study supported that ATP6V1B2 plays a vital role in normal hearing.
Beyond Hearing Sensitivity: Added value from Primary ABR Phenotypic Screening

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Background
The Sanger Institute's Mouse Genetics Project (MGP) generates targeted knockout mice and characterizes the effect of the mutant allele using a wide range of phenotyping tests designed to cover a number of common disease areas. Hearing is assessed in mice aged 14 weeks.

Methods
Mice are anaesthetized using Ketamine / Xylazine and their auditory brainstem responses (ABRs) recorded to click and a range of tone-pip stimuli. Thresholds are estimated and used to build audiometric profiles for mutant and control mice. In addition, click-evoked ABR waveforms are analysed in more detail. Individual responses recorded at 20dB and 50dB sensation level and a mean response waveform for each mutant group are superimposed on a reference range waveform calculated from large numbers of control wildtype mice of the same genetic background. These plots are used to assess altered waveform shapes in mutant mouse lines. Input-Output Functions are calculated for wave 1 and wave 3 amplitude and latency and plotted over a reference range for each parameter to quantify abnormal waveforms parameters.

Results
We have identified a number of mutant lines which demonstrated abnormal click-evoked ABR waveforms despite having normal threshold sensitivities. A variety of mutant lines were identified having affected wave 1 or wave 3 amplitude or latency. For example, Ppp5c homozygote mutants exhibited increased wave 1 amplitudes, Usp42 homozygote mutants had reduced wave 1 and wave 3 amplitudes, Atp5a1 heterozygote mutants showed reduced wave 3 amplitudes and Sesn3 homozygote mutants showed prolonged wave 3 latencies. Reduced wave amplitude could have a number of underlying causes. Reduced neuronal numbers could give reduced evoked potential generator activity and maybe indicate neurodegeneration phenotypes. Desynchronized neuronal activity could also result in reduced wave amplitude and may indicate abnormal synaptic activity along the auditory brainstem pathway. Altered wave latencies are also seen and may indicate deficits in neuronal conduction or synaptic efficiency.

Conclusion
Estimation of hearing sensitivity using ABRs can identify abnormal auditory phenotypes involving raised thresholds. However, detailed assessment of ABR waveforms can identify a wider range of potential auditory system phenotypes.

Collateral Damage: Spontaneous Mutations from a Targeted Knockout Programme

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Background
The Sanger Institute's Mouse Genetics Project (MGP) aims to generate knockout mice and characterise the effect of the null allele using a broad spectrum of phenotyping tests, one of which is the Auditory Brainstem Response. During standard phenotyping, several lines were found to display a variety of hearing defects which did not segregate with the knockout allele, and were probably due to spontaneous mutations.

Methods
DNA from four of these lines was sent for comparative genome hybridisation and whole exome sequencing, along with DNA from the embryonic stem cell lines used to make the knockout alleles. The mutations from two of the lines have also been mapped using a backcross. Further characterisation has been carried out by ABR, endocochlear potential (EP) measurements, middle ear dissection, inner ear clearing, immunohistochemistry and scanning electron microscopy of the organ of Corti.

Results
The sequencing results show a background of mutations present in ES cells, many common to both cell lines. Mapping has proven effective in identifying the mutation responsible for the rapid progressive hearing loss seen in one of the four lines, which has been identified as a new allele of S1pr2, a gene known to be important for hearing. These mice have normal gross morphology of middle and inner ears, and while hair cell degeneration was observed from 5 weeks old in affected mice, some intact hair cells are still visible at 9 weeks old in mice with no ABR responses at all. Their EP measurements, however, are decreased from an early age, and there are subtle defects in the stria vascularis of affected
mice. We are currently mapping a second line, and have identified a critical region and several candidates for the low frequency hearing loss phenotype common to this and several other MGP mouse lines.

**Conclusion**
The intriguing phenotypes of these spontaneous mutations are an unexpected benefit of the Mouse Genetics Project and its extensive phenotyping screen. The first mutation identified is a novel allele of a known hearing gene that offers a useful model of progressive hearing loss. Identifying the gene responsible for the unusual phenotype of low frequency hearing loss will further extend our understanding of the ear and the genes responsible for its maintenance and function.

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**Thrombospondin genes are required for normal cochlear development and maintenance**

**Diana Mendus**, felix wangsawihardja, **Vidya Sundaresan**, **Mirna Mustapha**

**Background**
Recent studies reveal important roles for cell-adhesion molecules such as thrombospondins (TSPs) in promoting synapse formation during brain development and in repairing synaptic connection after stroke.

**Methods**
For our efforts to identify genes involved in cochlear synapse formation and functional maturation, we hypothesize that cell adhesion proteins that are also upregulated during cochlear synaptogenesis and/or adulthood are important for these processes. In this study, we examine whether TSPs are involved in synapse formation and maintenance in the cochlea using recombinant mouse models of these genes.

**Results**
Our preliminary data using microarray and qPCR analysis on TSP genes 1-4 from whole cochlea revealed significant increase in TSP2-4 expression during a critical window of postnatal cochlear development with TSP1 expression picking up later in adulthood. Our auditory brain stem response (ABR) test on TSP1 and TSP2 mutants reveal elevated threshold in TSP2 mutants but not in TSP1 mutants at an early age. These data suggest a role for TSP2 in synapse functional maturation that is important for the onset of hearing. However, one-year old TSP1 mutant mice show an increase in ABR threshold at higher frequencies and decreased amplitude of response compared to wild type controls, suggesting a role for TSP1 in synapse maintenance. Our physiology data is in agreement with the roles predicated for these genes based on our expression studies. Counting of afferent and efferent synapse numbers in TSP2 and TSP1/2 mutants versus wild type mice does not show significant differences between these groups. Our next step would be to examine whether the synapses in the TSP mutant mice are functional using electrophysiology. To test the role for these genes in the recovery of the cochlea from noise injury, we exposed TSP2 and TSP1/2 mutant mice to a noise insult and measured ABR threshold after a recovery period. Data from these experiments showed that these mice were unable to recover from the noise injury compared to wild type controls indicating a deficit in repair mechanisms in these mice.

**Conclusion**
Our results, taken together, indicate that TSPs 1 and 2 are both involved in different aspects of cochlear functional maturation, repair and maintenance and these genes are now candidates for screening in human patients with congenital and age-related hearing impairment.
An Inducible rtTA Mouse Line for Gene Manipulation and Fate Mapping in Supporting Cells and Glia Within the Auditory and Balance Sense Organs of the Inner Ear

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Background
Sox10 is an SRY-related HMG-box (Sox) transcription factor that has been shown to be expressed in several cell types within the central and peripheral nervous systems. The reverse tetracycline transactivator (rtTA) protein is a variant of the E. coli tet-repressor protein that induces the expression of genes containing a tet-operator (tetO) sequence only when in the presence of tetracycline or its derivative doxycycline. Knock-in of the rtTA sequence to the endogenous Sox10 locus thereby allows for the specific expression of the rtTA protein in Sox10 expressing cells and temporal and quantitative control over TetO promoted gene expression by dosing regimen of doxycycline.

Methods
In this study, Sox10rtTA mice (MGI ID 3513104) were bred with mice containing a TetO-lacZ transgene, and X-gal staining and beta-galactosidase immunostaining were used to map Sox10 promoter activity in the sensory organs of the inner ear at various embryonic and postnatal ages. Sox10rtTA mice were also bred with mice that had a TetO-Cre transgene and a CAG-promoted, floxed-stop TdTomato reporter so that the fate of Sox10 positive cell types could be determined.

Results
Both models reveal widespread Sox10 reporter expression in the cochlea and balance organs at several ages and at varying levels across cell types depending upon age or upon the concentration of doxycycline administered. For both hearing and balance organs, the majority of reporter expression was in supporting cells (SCs) and glia, and, in the cochlea, in the greater epithelial ridge (GER). With late embryonic induction, lower doses of doxycycline resulted in decreased reporter expression in SCs, but not in GER cells, suggesting a dose-responsiveness for the TetO transgene, but perhaps also lower Sox10 promoter activity in SCs than GER cells. Fate mapping and reporter expression at embryonic and neonatal ages suggests that prosensory precursor cells that give rise to both HCs and SCs express Sox10, but that Sox10 expression is downregulated in HCs while being maintained in supporting cells well into adulthood.

Conclusion
These findings suggest that Sox10 may be an important factor in prosensory cell fate decisions and in the maintenance of SC fate in the hearing and balance organs. Furthermore, the Sox10rtTA mouse line represents a powerful model for fate mapping and/or genetic manipulation of Sox10 expressing SCs and glia in the mouse inner ear.

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The LINC Complex is Essential for Hearing in Humans and Mice
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Background
Young adults in two Israeli families of Iraqi Jewish ancestry presented with a high-frequency hearing impairment. The pattern of hearing loss in the families suggested autosomal recessive inheritance and the new deafness locus was named DFNB76. To obtain further insight into the role of Syne4 in epithelial cells, we created mice harboring a targeted disruption of the Syne4 gene by homologous recombination in embryonic stem cells. Hearing loss of mutant mice was detected already at P15 and progressed to all frequencies by P60. The Syne4\(^{-/}\) mice became a mouse model for DFNB76 once a SYNE4 mutation was found in the two families. Syne4 is a member of the KASH domain family. The nuclear envelope (NE), as the interface between the nucleus and cytoplasm, displays several major structural features, the most prominent of which are the inner and outer nuclear membranes (INM and ONM). Localization of KASH proteins to the ONM is dependent upon Sun domain proteins of the INM.

Methods
Linkage and homozygosity mapping, massively parallel and Sanger sequencing identified the human mutation. Mouse inner ears of Syne4 and Sun1 knock-outs were analyzed by scanning electron microscopy, auditory brainstem response (ABR), and behavioral testing. Immunolocalization of Syne4, Sun1, prestin and sodium potassium ATPase defined morphology and protein expression.

Results
SYNE4 was identified as the gene responsible for progressive high frequency hearing loss in the two families. Full-length Syne4 localized to the NE, whereas the mutant protein was found throughout the cytoplasm. In both Syne4 and Sun1 mutants, the OHCs began to degenerate in a basal to apical gradient, while the IHCs remained intact. Loss of Syne4 and Sun1 results in abnormal positioning of the nucleus in OHCs.

Conclusion
Hearing loss is caused by defects in either of two proteins that localize to the nuclear membranes of hair cells: Syne4, which localizes to the ONM and Sun1, which localizes to the INM. Mutations in SYNE4 lead to deafness in humans and the Syne4 mouse mutant serves as an ideal model to study this form of human deafness. The findings in the mice suggest that abnormal nuclear positioning leads to OHC death either due to interference with their sound-induced motility or to a compromised ability of abnormal OHCs to withstand the mechanical rigors of motility.

Supported by NIH/NIDCD grant R01-DC011835 and Singapore Biomedical Research Council and Singapore Agency for Science, Technology and Research (A*STAR).
Pathological Features in LmnaDhe/+ Mutant Mouse Provide a Novel Model of Human Otitis Media and Laminopathies

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Background
Genetic predisposition is recognized as an important pathogenesis factor of otitis media (OM) and associated diseases

Methods
The following assays were used:

Tympanometry analysis, ABR thresholds and DPOAE. histological and immunofluorescent staining. Scanning electronic microscopy, RT-PCR for gene expression. serum phosphate and calcium assays. Peritoneal macrophage migration in vivo and quantification assay

Results
We found that heterozygous mice with the Lmna allele, disheveled hair and ears (LmnaDhe/+), exhibit early-onset, profound hearing defects and other pathologic features of human laminopathy, which result from LMNA mutation. We assessed the effects of LmnaDhe/+ mutation on the development of OM and pathologic abnormalities characteristic of laminopathy. We found malformation and abnormal positioning of the Eustachian tube accompanied by OM in all of the LmnaDhe/+ mutant mice (100% penetrance) as early as postnatal day 12. Scanning electronic microscopy demonstrated ultrastructural damage to the cilia in middle ears that exhibited otitis media. Hearing assessment revealed significant hearing loss paralleling that in human OM. Expression of NF-κB, TNF-α and TGF-β, which correlated with inflammation and/or bony development, was upregulated in the ears or in the peritoneal macrophages of LmnaDhe/+ mutants. Rugous, disintegrative and enlarged nuclear morphology of peritoneal macrophages and hyperphosphatemia were found in LmnaDhe/+ mutant mice, and together these features resemble the pathology of human laminopathies possibly revealing some profound pathology, beyond OM, associated with the mutation.

Conclusion
The LmnaDhe/+ mutant mouse provides a novel model of human otitis media and laminopathy.
TRPM7 is required for neurotransmission by zebrafish hair cells
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Background
In an attempt to identify vertebrate genes required for hearing and balance, we undertook an examination of zebrafish mutants with defects in other forms of mechanosensation. One touch-unresponsive mutation, touchdown, arises in the gene encoding the transient-receptor-potential channel TRPM7. Because touchdown mutants also fail to initiate escape behaviors in response to water movements that are ordinarily detected by the lateral-line system, we attempted to identify the role of TRPM7 in hair cells of the lateral line.

Methods
We performed whole-mount in situ hybridization to pinpoint the cells in which TRPM7 is expressed. Electrophysiological recordings were then made from neuromast hair cells and lateral-line afferent neurons in touchdown mutants, wild-type siblings, and mutants in which the expression of wild-type TRPM7 had been selectively restored in hair cells. Finally, we examined the afferent synapses of hair cells by electron microscopy, immunohistochemistry, and pH-sensitive probes.

Results
Expression analysis demonstrated trpm7 transcripts in neuromast hair cells. Electrophysiological recordings from hair cells and their afferent neurons in touchdown mutants revealed normal microphonic responses but no tonic, hair cell-driven neural activity. Consistent with a requirement for the channel in hair cells, cell-specific expression of TRPM7 restored tonic activity in the afferent neurons of mutants. These findings, coupled with a role of mammalian TRPM7 in neurotransmission by cultured neurons, prompted a structural investigation of hair-cell synapses in the mutants. We found that touchdown larvae possess the structural elements of hair-cell synapses but lack acidified synaptic vesicles.

Conclusion
Our results demonstrate that TRPM7 is required for neurotransmission by zebrafish hair cells and suggest that the channel is necessary for proper pH regulation of synaptic vesicles, a process required for neurotransmitter filling.
Role of the ATP-gated P2X2 Ion Channel in Progressive Nonsyndromic Hearing Loss DFNA41 and in Noise-related and Age-related Hearing Loss

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Background
Extracellular ATP is a purinergic signaling molecule that influences developing and mature cochlea via both ionotropic P2X and metabotropic P2Y receptors in sensory, neural, and secretory cochlear tissues. A role of ATP signaling in regulating hearing sensitivity to noise is supported by evidence that noise and metabolic stress trigger ATP release into the endolymph. Sustained noise exposure also causes up-regulation of P2X2 receptor mRNA and protein expression in the organ of Corti, that is highly expressed in developing inner ear neuroepithelium. There is evidence in both mice and humans that there is a heritable component to noise-induced hearing loss, with no known genes directly associated with noise- and age-related human hearing loss. We identified P2RX2 p.V60L variation that co-segregates with inherited dominant and progressive hearing loss DFNA41, and determined how it affects the function of the P2X2 channel.

Methods
We used patch-clamping measurement of HEK293 cells expressing GFP-P2RX2 to evaluate its physiology. We used confocal microscopy of MDCKII cells expressing GFP-P2RX2 to quantify its ATP-induced permeability to FM1-43 dye. We performed ballistic transfections and confocal microscopy to examine the targeting properties of GFP-P2X2 in inner ear hair cells and support cells. We examined Auditory Brainstem Recordings and inner ear histology of aging mice and of mice exposed to noise from P2X2 (-/-) knockout colony.

Results
We show that both wild-type and p.V60L mutant P2X2 receptors are preferentially targeted to the apical plasma membranes of hair cells and supporting cells, excluding mislocalization of the heteromeric channel as a cause of deafness in heterozygous individuals. We also show that P2X2 p.V60L receptors have aberrant electrophysiological characteristics, and that co-expression of mutant and wild-type subunits results in severe reduction (~60%) of permeability of the heteromeric channel to FM1-43 dye, suggesting a dominant-negative effect of the mutant protein on channel function. Moreover, stimulation by ATP evokes an inward current response in cells expressing wild-type P2RX2, but not in cells expressing P2RX2 p.V60L. Finally, we find that loss of function of P2X2 receptors in mouse contributes to noise-related and age-related hearing loss.

Conclusion
Critical functions of P2X2 receptors are abolished by the P2RX2 p.V60L mutation. These results indicate that P2RX2 p.V60L is responsible for DFNA41 and that loss of function of P2X2 receptors contributes to both noise-related and age-related hearing loss.

Defining the Pendred Syndrome-DFNB4 Disease Spectrum by High Throughput Sequencing of 101 Candidate Genes Expressed in the Lateral Wall of the Murine Membranous Labyrinthine in Patients with Enlarged Vestibular Aqueducts

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Background
Pendred syndrome (PS) and non-syndromic enlarged vestibular aqueduct (EVA) are recessive disorders characterized by sensorineural hearing loss and inner ear malformations that range from Mondini dysplasia to isolated EVA. Mutations in SLC26A4, encoding the anion exchanger pendrin, are the major cause of the PS-EVA-DFNB4 disease spectrum. However, in a small percentage of patients carrying only a single mutation in SLC26A4, additional mutations have been
identified in cis- and trans- regulatory elements of SLC26A4 (its promoter and the transcriptional regulator FOXI1), as well as in the potassium channel KCNJ10.

Methods
To identify additional genes that may interact with pendrin, we have used a microarray-based strategy to quantitate gene expression in the lateral cochlear wall in mice with a targeted deletion of Slc26a4 (Slc26a4-/- mutants). To select candidate genes for further study, network analysis was used to define gene-gene correlations based on expression data, filtering for all possible interacting partners of SLC26A4, FOXI1 and KCNJ10. We also included genes implicated in murine deafness for which no human counterpart has been identified (Heatmap database).

Results
Targeted genomic enrichment has been used to capture all coding sequence of genes on our candidate list. The genes included on the optimized platform, which we call EVASeq, are composed of potassium channels, sodium channels, solute carriers, carbonic anhydrases, and some genes related to ATP catalysis. In addition, we have included 3 known EVA genes, 4 known deafness genes and some genes with other functions. To comprehensively define the EVA phenotype, all patients with EVA and one or no mutations in SLC26A4 are being screened for variants in both EVASeq genes and OtoSCOPE genes. Data analysis is performed using a locally installed Galaxy framework running a custom pipeline for variation annotation.

Conclusion
The results of this study will provide novel insight into the PS-EVA-DFNB4 disease spectrum.
<table>
<thead>
<tr>
<th>Targeted gene categories included on EVASeq</th>
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<tbody>
<tr>
<td>Potassium Channel genes</td>
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<tr>
<td>Sodium Channel genes</td>
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<tr>
<td>Solute Carrier genes</td>
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<tr>
<td>Carbonic Anhydrase genes</td>
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<tr>
<td>Other Trans-membrane genes</td>
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<td>ATP Catalyzes</td>
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<tr>
<td>Known EVA genes</td>
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<td>Known deafness genes</td>
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<td>Others</td>
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**Mutation of SLC45A3 in familial and sporadic Menière’s disease**

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**Background**

Menière’s disease (MD) is a complex idiopathic disorder of the inner ear characterized by the symptom triad of vertigo, sensorineural hearing loss and tinnitus. Its incidence varies around the world and is estimated at 1-2 per 1,000 Caucasian individuals. Most cases of MD are sporadic although 5-14% of individuals report a family history in which MD segregates as a dominant but incompletely penetrant disease. To date, a genetic contribution to sporadic and/or familial MD has not been identified reflecting the challenges that complex diseases present to causative gene discovery.

**Methods**

We completed linkage analysis in a large family segregating MD and confirmed genetic findings in 570 patients with sporadic MD. Expression of the gene and its encoded protein in the membranous labyrinth were defined by in situ hybridization and immunohistochemistry.

**Results**

The genome-wide linkage scan defined a candidate interval on 1q32 that segregated with the MD phenotype. Targeted exon capture and pyrosequencing of coding sequence within the interval identified a 36-bp deletion in SLC45A3. We found the identical deletion in five of 570 patients with sporadic MD. In situ hybridization and immunohistochemistry confirmed that SLC45A3 and its encoded protein, Prostein, are expressed in the cochlea and endolymphatic sac.

**Conclusion**

Through the use of two complementary human genetics approaches – 1) segregation analysis with targeted-sequence capture and massively parallel sequencing; and, 2) candidate gene sequencing in a large MD cohort as a test of the common-disease-rare-variant hypothesis – we have identified SLC45A3 as the first gene causally related to MD. A deletion in SLC45A3 is found in 1% of persons with MD across five populations. The encoded protein, Prostein is expressed in the cochlea and endolymphatic sac where it is hypothesized to play a role in osmotic regulation. (This research was supported in part by a grant from the American Otological Society (RJHS).)
The Static Force Dependence of Sound - Transmission Efficiency in Round Window Stimulation
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Background
The Direct Acoustic Cochlea Stimulator (DACS, Phonak Acoustic Implants) is intended to stimulate the cochlea directly by a piston. An alternative approach to the round window (RW) is successfully done with other devices, having the advantage of being independent of the presence of any middle ear structure. Here we investigate the impact of static force applied to the RW on actuator output and performance.

Methods
The obtainable maximum equivalent sound pressure output of round window stimulation was determined experimentally in fresh human temporal bones (TBs) in analogy to the ASTM standard (F2504.24930-1).

First the stapes footplate (SFP) displacement in response to sound applied to the external ear canal was determined. ASTM compliant TBs were implanted with the actuator axis perpendicular to the RW membrane. The stimulation was performed by means of a prosthesis with a spherical tip (Ø 0.5mm) attached to the actuator as interface to the RW membrane.

SFP vibration in response to actuator stimulation was measured in a series of positions relative to the RW membrane. Starting at a position without contact to the RW, the position of the actuator was move in increments of 50 μm into the RW membrane and the resulting axial static force load was determined. From the SFP response to sound and actuator stimulation the achieved equivalent sound pressure level at the tympanic membrane was calculated for each step.

Results
10 TBs were found within the acceptance range of the ASTM standard and contributed data to further analysis. At moderate axial static force (3.90 ± 0.70 mN; MV, SD) an average sound equivalent sound pressure level of ~103-120 eq. dB SPL (@1Vrms) was found for frequencies ≤ 4 kHz. At higher frequencies (6 - 10 kHz) the achieved output dropped to ~90 dB SPL. This static force loading was below the actuator safe operating limit (< 50 mN) and at least a factor 10 below the rupture limit of the RW.

The magnitude of the transfer function from the RW to the SFP increased monotonically with the applied forces (0 to >40 mN), allowing even higher output by increasing the axial load.

Conclusion
Sound transmission efficiency to the RW crucially depends on the static contact force applied to the RW. Our results demonstrate the possibility of round window stimulation with the DACS actuator at moderate static forces with sufficient output to treat moderate and pronounced sensor neural hearing loss components.

Estimating round window vibroplasty efficiency: A comparative temporal bone study of forward and reverse ossicular chain stimulation
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Background
Mechanical stimulation of the round window membrane (round window vibroplasty, RWV) with an active middle ear implant (AMEI) has demonstrated functional benefit in clinical reports in patients with mixed and conductive hearing losses. Current published standards for objective measurement of AMEI performance are based on stapes velocity in human cadaver temporal bones for AMEIs coupled to the ossicular chain (ossicular chain vibroplasty, OCV). Here, stapes velocity is a reasonable measure of the input to the cochlea via the AMEI because the system is driven in the forward direction. However, these standards may not be applicable when estimating the efficiency of RWV as the stapes or umbo velocities may change in complex ways though stimulation of the cochlea by the round window. Previous work from our lab in an animal model revealed stapes velocity measures underestimate the RWV efficiency for frequencies above 4 kHz compared to cochlear microphonic measurements. In the current study, we sought to expand upon this hypothesis.

Methods
In the current set of temporal bone experiments (N=5), we measured stapes and umbo velocities, Hsv and Husv respectively, via laser Doppler vibrometry in response to acoustic and AMEI generated tone pip stimuli (0.25 to 8 kHz)
delivered by OCV and RWV for varying intensity levels. $H_{sv}$ and $H_{uv}$ in RWV and OCV were normalized with the representative acoustic $H_{sv}$ and $H_{uv}$ to estimate the AMEI efficiency in each condition, which was analyzed in 3 frequency ranges (low 0.25-1 kHz, med 1-3 kHz and high 3-8 kHz).

**Results**
In comparing the efficiency estimated from $H_{sv}$ and $H_{uv}$ in RWV, measurements revealed that the $H_{uv}$ underestimated AMEI efficiency by 10, 6 and 15 dB (low, medium and high frequency ranges, respectively). In OCV, $H_{uv}$ underestimated AMEI efficiency by 5 dB, 13 dB and 21 dB (low, medium and high frequency ranges, respectively).

**Conclusion**
The vibromechanical energy delivered by RWV measured at the stapes or umbo is subjected to losses from the cochlea and the ossicular chain. Transmission losses are proposed to occur secondary to third window effects within the cochlea in addition to ossicular chain inefficiencies. With the addition of intracochlear pressure measures, better estimation of efficiency of RWV may be obtained in future studies.

**Use of Temporal Envelope Cues Recovered from Temporal Fine Structure for Cochlear Implant Users When Original Speech Envelopes Are Severely Degraded**

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**Background**
Won et al. [J. Acoust. Soc. Am. 132, 1113-1119 (2012)] reported that cochlear implant (CI) sound processor generates recovered amplitude modulation (AM) cues from broad-band speech frequency modulation (FM) and CI users can use these cues for speech recognition in quiet. For commercially available implant processing today, such FM-to-AM conversion is the only mechanism of transmitting FM (i.e., acoustic temporal fine structure, TFS) information for CIs. Therefore it is important to further study the role of FM-to-AM conversion for CI users in various acoustic environments. The present study aimed to determine whether the FM-to-AM conversion mechanism is important for CI users when original speech AM cues are severely degraded.

**Methods**
Original speech AM cues were degraded by either manipulation of the acoustic signals or of the sound processor. The manipulation of the acoustic signals included the presentation of background noise, simulation of reverberation, and amplitude compression. The manipulation of the sound processor included the change of input dynamic range and the number of channels. For each of these conditions, multiple levels of envelope degradation were tested. Speech perception was measured for CI users and compared for stimuli having both AM and FM information (intact condition) or FM information alone (FM condition). Consonant identification or word recognition were used as speech perception measures.

**Results**
As original speech envelopes were degraded more, speech perception performance was reduced for both intact and FM conditions, but performance was always better than chance. Performance for intact and FM conditions became similar for stimuli having severe envelope degradation, suggesting that speech perception for CI users in such a challenging acoustic environment could be determined by the use of recovered envelope cues. This is further supported by significant correlations between speech perception for intact and FM stimuli as well as electrode output measurements.

**Conclusion**
The present study demonstrates that the envelope cues recovered from speech TFS convey important phonetic information for CI users when they are communicating in an acoustic environment which degrades the original speech envelope cues.

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Efficient coding for auditory prostheses
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Background
Conventional cochlear implant (CI) sound coding strategies (e.g. CIS, ACE) convey information from multiple spectral bands by sampling the temporal envelope of each channel with fixed-rate pulses. We investigated an experimental sound coding strategy, called Fundamental Asynchronous Stimulus Timing (FAST), which employs a sparse, yet temporally precise, representation of temporal envelope. We hypothesized that the FAST strategy would provide better temporal sensitivity to ITD and F0 cues while using significantly less stimulation power than conventional strategies.

Methods
Several types of listening experiments were conducted comparing the ACE and FAST strategies in bilateral and unilateral CI subjects. These include 1) speech understanding in quiet and in noise, 2) ITD sensitivity of speech stimuli, 3) ITD-based spatial release from masking, and 4) F0 sensitivity. Additionally, battery life during take-home usage was examined to compare power consumption across strategies.

Results
Initial results indicate that speech understanding is equivalent across strategies, while FAST provides significantly better ITD sensitivity and ITD-based spatial release from masking. Additionally, subjects reported 2-3 times more battery life with FAST over ACE.

Conclusion
Results suggest that a temporally sparse coding strategy, such as FAST, provides a sufficient representation of speech sounds while allowing for more precise and efficient coding of temporal envelope. This development could lead to better hearing outcomes with CIs and future device miniaturization.
Benefits of Intermodal Training for Sound Localization following Bilateral Cochlear Implantation

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Background
Arguably, cochlear implants (CIs) are the most successful neural prosthesis developed. Although most individuals are fitted with a CI in one ear only, bilateral CIs are increasingly being offered to children with the aim of improving sound localization and speech perception in noise. However, adults with long-term hearing loss are rarely offered bilateral CIs. We developed an animal model of bilateral CI (Hartley et al., 2010) to examine the potential detrimental effects of long-term hearing loss on sound localization following bilateral CI and possible strategies to remediate them.

Methods
Ferrets were fitted with bilateral (n=4) or unilateral (n=4) CIs after either early (n=4) or late (n=4) induction of hearing loss by ototoxic antibiotic injections. They were subsequently assessed for their sound localization accuracy within a free-field environment, using a positive conditioning paradigm. Animals that performed poorly were reassessed after a modified intermodal training paradigm that consisted of localization trials in which visual cues (light flashes) were randomly interleaved with sound-alone stimuli. In acute physiological recordings that followed behavioral assessments, we examined interaural timing difference (ITD) and interaural level difference (ILD) sensitivity of primary auditory cortical (A1) neurons in response to single biphasic current pulses presented binaurally to the intracochlear electrode arrays under direct, computer control.

Results
Compared with normal-hearing ferrets, animals with late-onset hearing loss and bilateral CI performed poorly on the sound localization task. However, these animals performed significantly better than bilaterally implanted ferrets with early-onset hearing loss or animals with unilateral CI. The latter performed no better than chance, regardless of the duration of deafness. Strikingly, intermodal training led to significantly improved sound localization accuracy in bilaterally implanted ferrets with early-onset hearing loss. In these animals, cortical sensitivity to ITDs and ILDs in A1 was much greater than in ferrets that did not receive intermodal training.

Conclusion
Our results show that despite the absence of auditory input during development, intermodal training led to improved sound localization accuracy in animals with bilateral CIs. Our recording data reveal a possible neural substrate for this by showing improved binaural sensitivity of A1 neurons in these animals. These findings support investigation of a similar paradigm for auditory rehabilitation in human CI users, specifically where current implantation criteria exclude bilateral implantation in individuals with pre-lingual onset of hearing loss.

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Early deafened, Late-implanted Cochlear-Implant users enjoy Listening to Music
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Background
The early-deafened, late implanted (EDLI) CI users are an only recently implanted, and therefore understudied, group of CI users. The outcomes of implantation are mostly unknown for this group. Therefore, this study evaluated the outcome of implantation in EDLI CI users for the second most important acoustical stimulus next to speech, the self-perceived perception and enjoyment of music. Additionally, the correlations of these measures were explored with the self-reported quality of life, the self-reported everyday hearing ability and a behaviorally measured speech perception test.

Methods
The inclusion criteria for participation were: severe hearing loss at least since preschool (5-7 years of age), implanted > 16 years and > one year of CI-experience. Thirty-seven qualifying EDLI CI users, all patients of our clinic, were sent four questionnaires to quantify: 1) Dutch Musical Background Questionnaire (enjoyment and perception of music); 2) Nijmegen Cochlear Implant Questionnaire (health-related quality of life); 3) Cochlear Implant Functioning Index (auditory related functioning); 4) Speech, Spatial and Qualities of Hearing scale (hearing ability). As a complementary behavioral measure, a speech perception in quiet test (phoneme recognition in meaningful words reported in percent correct) was completed.

Results
Twenty-two EDLI CI users were included. A majority (60%; 12 out of 22) reported music to sound pleasant through their CI. The subjective quality of the sound of music was scored positively. No correlations were observed between the self-reported perception and enjoyment of music, the quality of life, everyday hearing ability and the behavioral scores of speech perception. The self-reported health-related quality of life of the NCIQ and the auditory related functioning of the CIFI were positively correlated with the behavioral measures of speech perception.

Conclusion
The overall results indicate that, differently than for post-lingually deafened CI users, the group of EDLI CI users subjectively enjoy the perception of music and rate the quality of music positively. No correlations between the subjective perception of music and the health-related quality of life, everyday hearing ability and speech perception were shown. An explanation for the absence of correlations could be that this group of CI users rates their perception of music to a different anchor point than the post-lingually deafened adults, which may be attributed to the lack of exposure to music with acoustical hearing. The findings of our study may give additional support for implantation of this special group.

Musical Interval Perception with Normal Hearing and Cochlear Implants
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Background
Little is known about cochlear implant (CI) users’ ability to perceive pitch intervals (i.e., the size of pitch changes between notes) important for melody recognition. This study systemically investigated musical interval perception of CI users via clinical processors using both subjective and objective tests that are suitable for patients even with little musical experience.

Methods
Subjects were adult CI users (n=6) and normal-hearing (NH) listeners (n=6). Interval size estimation was tested for musical intervals from 1 to 24 semitones using a scale from 1 to 100. Pitch interval discrimination threshold was measured for standard intervals from 0 to 9 semitones using a 2AFC task and an adaptive procedure. Both rising and falling intervals were tested in a high (around 349.2Hz) or a low (around 174.6Hz) pitch range. In the melodic interval adjustment test, subjects adjusted a scaling factor that started from a random value and was applied to all the intervals of a familiar melody to produce the most correct melody. Subjects were also asked to rate the degree that each adjusted melody was in tune.

Results
CI users had poorer performance than NH listeners. Estimated interval sizes were generally a linear increasing function of physical interval sizes in semitones for NH listeners, but a stepwise increasing function for CI users. Estimated sizes increased more rapidly for falling intervals than for rising intervals in the low pitch range, but for rising intervals than for falling intervals in the high pitch range. The pitch interval discrimination thresholds increased as the standard interval
increased from 0 to 1 semitone and from 7 to 9 semitones, but were relatively constant for standard intervals from 1 to 7 semitones. There were no consistent patterns in interval discrimination thresholds between falling and rising intervals in the low and high pitch ranges. In the melodic interval adjustment test, NH listeners were sensitive to subtle interval changes and found the melodies with an interval scaling factor of ~1 the most correct. CI users often preferred expanded intervals with scaling factors >1. However, the found scaling factors varied across trials, melodies, and subjects, and seldom produced a completely in-tune melody.

**Conclusion**
The results revealed significant effects of pitch range and direction on musical interval perception. Subjective estimated interval sizes cannot be explained by objective interval discrimination thresholds. Subjects with better interval discrimination seemed to have better judgments of musical intervals in familiar melodies.

36 Vocal Singing in Prelingually-Deafened Children with Cochlear Implants and Hearing Aids
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**Background**
Vocal singing plays an important role in people's social and communicative aspects of daily life. Profoundly deaf children who use cochlear implants or hearing aids frequently have difficulty perceiving and producing discrete pitch patterns. The purpose of the present study is to investigate vocal singing abilities in children with cochlear implants and hearing aids.

**Methods**
Three groups of participants, aged between 2 and 7 years old, were recruited: (1) nine cochlear implant users, (2) twelve hearing aid users, and (3) ten normal-hearing, age-match children. Each child was asked to sing a familiar song. The fundamental frequency (F0) and duration of each note of the recorded songs was extracted. The sung songs were then compared against the original score based on the following five different metrics: (1) contour direction, (2) compression ratio, (3) mean note deviation, (4) mean interval deviation, and (5) note duration ratio deviation. The first four metrics were pitch-based measures and the fifth one was a rhythm-based measure.

**Results**
Results showed that children who use cochlear implants and hearing aids performed similarly with each other in pitch- and rhythm-based measures, but the performance was significantly poorer than that of the normal-hearing children.

**Conclusion**
Either cochlear implant or hearing aid has failed to provide satisfactory rehabilitation to prelingually-deafened children in vocal singing due to inadequate pitch information that is transmitted to the auditory system through the devices.

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37 Is high intensity electrical stimulation excitotoxic in hearing cochleae? Evidence from the mouse model.
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**Background**
Excitotoxicity after high intensity electrical stimulation is one proposed mechanism for hearing loss seen after cochlear implantation in patients with residual hearing. Consistent with this hypothesis, data from cochleotypic cultures exposed to high intensity electrical current reveal damage at the level of the primary afferent synapse with relative sparing of hair cells.

**Methods**
33 three month old C57Bl6J mice underwent round window implantation with a platinum wire electrode. 20 of these were exposed to acute electrical stimulation (ES: 100Hz monopolar charge balanced biphasic pulses) at high intensity (eABR saturation). Hearing was tested using ABR and DPOAE pre-implantation and at 1 week. After euthanasia (1 week),
cochleae were prepared for either histopathology (5 ES, 4 sham) or immunohistochemistry (15 ES, 9 sham). Contralateral ears were also examined.

Results
Post-implantation ABR thresholds were significantly elevated in sham and electrically stimulated cochleae compared with pre-implantation and contralateral thresholds with a trend towards higher thresholds in electrically stimulated cochleae. DPOAEs were undetectable in cochleae with the highest thresholds. Histology revealed hair cell loss in 2/5 stimulated and 0/4 unstimulated mice. Interestingly, if hair cell loss was evident in the basal turn – it was also evident in the mid cochlea. Currently we are quantifying hair cells, afferent synaptic structures, and neural innervation based on immunolabeling.

Conclusion
C57Bl6J mice suffer a large decrement in hearing (42+/−4dB) after round window implantation – likely due, in part, to the proportionate effect of middle ear effusion relative to size. Loss of hair cells is apparent only in stimulated mice. These data provide support for an “electrotoxic effect” but suggest that only mice lacking significant middle ear effusion (intact DPOAEs) after implantation should be used in future analyses. Work is ongoing to pinpoint the cochlear structures most affected.

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Effects of Age on the Preservation of Residual Hearing with Cochlear Implants
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Background
Cochlear implant (CI) recipients with residual hearing in their implanted ear receive both electric and acoustic stimulation. As a result, optimal hearing preservation is a major factor in achieving the best possible auditory experience for these patients. However, a small but significant number of CI patients lose their residual hearing following implantation, although the mechanisms of this loss are not clear.

Methods
To determine the effects of age on the preservation of residual hearing, this study evaluated hearing preservation, using auditory brainstem response recordings, in both neonatal (n=8) and adult onset (n=3) partially deaf cats. All animals received approximately 6 months intracochlear electrical stimulation from a clinical CI and speech processor, after stabilisation of their systemic aminoglycoside induced partial hearing loss.

Results
All animals had measurable residual hearing in the low- to mid-frequency range (below 8 kHz, apical to the intra-cochlear electrode array) throughout the experiment. There was no significant loss of hearing in the neonatally deafened animals during the chronic stimulation period (paired T-test, p= 0.13). In contrast, the adult onset group exhibited a significant hearing loss of approximately 8 dB (paired T-test, p= 0.03). Interestingly, there was no difference in the severity of the loss between the low- (1 & 2 kHz) and mid-frequency (4 & 8 kHz) regions (t-Test, p = 0.5).

Conclusion
These results indicate that CI use per se, does not result in a significant loss of residual hearing (i.e. no change in threshold for the neonatally deafened group). However, age may be a factor in the preservation of residual hearing with CI use, as a small loss was observed in the adult onset group. These results need to be confirmed with a larger sample size, and confounding factors (such as the total duration of deafness) also need to be further examined.

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The effects of neurotrophins and chronic electrical stimulation delivered to the deafened guinea pig cochlea
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Background
Spiral ganglion neurons (SGNs) in the deafened cochlea undergo continual degeneration ultimately resulting in cell death. The exogenous application of neurotrophins (NTs) can prevent SGN loss, with the survival effects enhanced by chronic intracochlear electrical stimulation (ES) from a cochlear implant. Furthermore, NT treatment can enhance resprouting of the SGN peripheral processes. However, little is known about the changes to the SGNs that occur following ES and NT
treatment. Here we examine the ultrastructural changes to SGNs and their peripheral processes following delivery of brain derived neurotrophic factor (BDNF) and chronic ES.

Methods
Adult guinea pigs (n=14) were deafened using a single application of an aminoglycoside and loop-diuretic. Two weeks later each animal was unilaterally implanted with a scala tympani electrode array containing a cannula for NT delivery. A commercial cochlear implant was used to deliver chronic intracochlear ES over a four week treatment period. Cochleae were collected and prepared for examination on a transmission electron microscope.

Results
Chronic NT treatment was effective in reducing the loss of SGNs and their peripheral processes following deafness. SGN soma in deafened controls exhibited enlarged cytoplasmic vacuoles while the myelin sheath typically remained intact. The peripheral processes were significantly larger in NT treated cochleae, with or without ES, compared to deafened cochleae not treated with NTs (p<0.0005). Finally, resprouting processes were observed within the osseous spiral lamina, the spiral limbus and the scala tympani.

Conclusion
This study has shown that exogenous NT delivery was effective in reducing the retraction of peripheral processes and SGN degeneration that normally occurs following deafness. Degeneration of the SGN soma precedes loss of the myelin sheath. Resprouting of peripheral processes was promoted by NT treatment. Processes were observed within the scala tympani, raising the potential of a direct connection between the SGNs and the electrode array that may improve the nerve-electrode interface.

Safety of Intratympanic Gel Polymer Injection for Treatment of Cytomegalovirus-Induced Hearing Loss
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Background
Cytomegalovirus (CMV) is the leading infectious cause of congenital hearing loss, affecting 0.6 per 1000 newborns each year. Intratympanic injection of cidofovir (CDV) can rescue hearing in CMV-infected guinea pigs, but therapeutic benefit is limited by drug loss through the Eustachian tube. Sustained delivery of CDV to the intracochlear space can be accomplished by direct application of drug-infused gel polymer to the round window membrane. However, the safety of this technique is poorly described. The objective of this study is to determine whether intratympanic delivery of CDV using a gel polymer is a safe treatment for CMV-induced hearing loss.

Methods
Non-infected guinea pigs were injected intratympanically with one of four gel polymer combinations (Gel A, Gel B, Gel A + CDV, Gel B + CDV). Auditory brainstem response measurements were obtained prior to injection and at 21, 28, or 35 days post-injection. Hearing loss was classified as mild (<20 dB), moderate (20-40 dB), or profound (>40 dB) based on decibels lost in post-treatment hearing thresholds. The middle ear and cochlea were then evaluated microscopically for signs of inflammation (edema, cell infiltrate, bleeding). Inflammation was classified as mild (1/3), moderate (2/3), or severe (3/3) based on the number of inflammatory factors present.

Results
Moderate-to-profound high frequency hearing loss and moderate-to-severe ear inflammation were measured in all treated guinea pigs regardless of gel polymer combination used. Hearing loss and inflammation were maximal at day 21 and improved by day 35. However, measurements never returned to pre-treatment baseline.

Conclusion
Intratympanic injection of CDV-infused gel polymer holds great potential as treatment for CMV-induced hearing loss, but has not yet reached an acceptable level of safety. Further research is warranted to develop a safe method of drug delivery that may one day provide treatment for affected infants.
Protein Transduction utilizing Arginine-rich Cell-Penetrating Peptides into the Cochlea.

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Background
Various approaches, including viral vectors, electroporation, microinjection, and liposome encapsulation, have been used to introduce target molecules into the cells to manipulate them. In 1988, Green and Frankel independently discovered that HIV TAT proteins were able to cross cell membranes. Short peptide sequences, such as these, have been referred to as cell-penetrating peptides (CPPs). CPPs are a class of short cationic peptides that are able to traverse the cell membranes in many types of mammalian cells. Many macromolecules have been attached to these peptides and subsequently internalized. The cargo molecules delivered into cells maintain their biological activities. Among CPPs, arginine-rich CPPs have been the most widely studied. We assessed protein transducing abilities of nine-arginine-rich CPP into embryonic inner ears and adult inner ears.

Methods
Ptoteintransduction into the developing inner ear: CD-1 pregnant mouse dams with E11.5 embryos were anesthetized. The uterus was exposed using a low midline laparotomy. EGFP conjugated to nine-arginine peptide (EGFP-9R), or EGFP alone was injected into the otocysts.
Ptoteintransduction into the adult inner ear: Gelatin foam with soaked in EGFP-9R or EGFP alone in was placed at the round window of adult CD-1 mouse cochleae.
Protein transducing abilities was assessed periodically.

Results
In the embryos that underwent EGFP-9R inoculation, EGFP-9R expression was detected in the lining cells in the otocysts and in their vicinity in a diffuse manner. Expression of EGFP-9R was maintained for 12-18 hours after otocystic inoculation. Embryos that underwent EGFP-9R inoculation developed to term normally, and P30 mice that underwent EGFP-9R inoculation showed normal auditory function.
In adult EGFP-9R mice, EGFP-9R expression was detectable in the cocholeautp to 36 hours post-treatment.

Conclusion
Ptoteintransduction may provide a useful strategy for delivering a target molecules into embryonic inner ear and/or adult mice inner ear. Ptoteintransduction via the round window membrane may be a potentially useful method for delivering therapeutically relevant molecules into the inner ear.

Steroid Treatment Partially Suppresses the Altered Inner Ear Gene Expression in Murine Autoimmune Disease

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Background
It has been decades since the first identification of inner ear disorders in systemic autoimmune diseases and their control by glucocorticoids. However, little is known of the mechanisms by which systemic inflammatory diseases cause hearing loss or the molecular impact of steroids on these dysfunctional processes. Although hearing recovery is generally attributed to the immune suppressive functions of glucocorticoids, they also are known to affect inner ear mineralocorticoid-receptor driven processes of ion and water homeostasis. Clarification of these steroid controlled mechanisms in the ear is critical for effective therapeutic control of many forms of hearing loss. Therefore, studies were conducted to assess the inner ear molecular processes at risk in murine lupus and the impact of glucocorticoid treatment.

Methods
Systemic disease and hearing loss develop in autoimmune MRL/MpJ-Fas/J mice at 3 months of age. Therefore, inner ear gene analyses were conducted on 3.5 week old and 5.5 month old mice, as well as 5.5 month old mice treated systemically with the glucocorticoid prednisolone for 2 weeks. Inner ears were harvested, RNA extracted, and 6 ear samples for each treatment group processed on the Affymetrix 430 2.0 Gene Chip. Genes for the treated and untreated older mice were statistically analyzed for up or down expression against the young mice.

Results
Approximately 7,500 genes were significantly up or down regulated by systemic disease (FDR p < 0.05). If a cutoff of 1.5 fold or more overexpression or 0.67 or less underexpression was applied to these significantly affected genes, there still
were 1,303 (up) and 1,029 (down) inner ear genes affected. Steroid treatment reduced this number by approximately 20%. Inflammation related genes and ion homeostasis related genes were examined further since these two mechanisms presumably play a significant role in cochlear function during autoimmune disease. Steroid treatment reduced the number of overexpressed inflammatory genes by 30% and decreased the number of underexpressed ion homeostasis genes by 50%.

**Conclusion**

Thousands of inner ear genes are affected by systemic autoimmune disease and oral steroids play a role by reversing some of these induced molecular processes. Furthermore, the efficacy of steroids in controlling hearing loss probably involves more inner ear mechanisms than simple immune suppression.

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**Magnetic nanoparticles to deliver drugs to the ear**

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**Background**

In the inner ear, noise trauma can induce injury and inflammation. The damage should be ideally treated by the timely local delivery of drugs to the affected relevant structures in a limited region of the cochlea. To this end, nanoparticles are a natural candidate because of their biocompatibility, the ability to load them with a variety of drugs, and the promise that they can deliver their payloads without causing additional injury or trauma to the inner ear.

**Methods**

We are using superparamagnetic nanoparticles with a maghemite core, made biocompatible with a chitosan matrix loaded with fluorescent proteins for visualization. We have characterized the distribution of these particles in the cochlear space as a function of their diameter, duration of exposure to an external magnetic field and other delivery parameters while the particles are being actively steered by a configuration of magnets, either pulling from the contralateral side of the skull or pushing from the ipsilateral side.

**Results**

We are able to deliver nanoparticles with a remarkably uniform density throughout a target area of the cochlea, thus leading to the hope we can deliver drugs in a targeted, uniform manner at a pre-determined therapeutic level. We have also tested in a rat model the effectiveness of magnetically pushed nanoparticles functionalized with prednisolone in preventing the emergence of hearing loss post-trauma.

Finally, we have also observed the possibility of using the nanoparticles to deliver drugs from the ear canal into the middle ear, with a view to cure or reduce otitis media without the need to administer drugs systemically or to compromise the integrity of the eardrum.

**Conclusion**

The use of biocompatible magnetic nanoparticles, functionalized with appropriate molecular agents, will lead to the ability to deliver specific drugs at pre-determined concentrations in specific parts of the ear.
NIDCD Workshops: Training/Career Development Workshop and Early Stage/New Investigator Workshop

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¹NIDCD

Background

NIDCD will offer two concurrent workshops targeted to two specific audiences with limited experience submitting grant applications to NIH.

Methods

Workshop #1: Applying for NIDCD Training and Career Development Grants.

Workshop #2: Early Stage Investigators (ESI) and New Investigators (NI).

Results

Workshop #1: Applying for NIDCD Training and Career Development Grants.

This workshop will include an overview of research training and career development opportunities appropriate for graduate students, postdoctoral fellows and new clinician investigators. The discussion will include the submission and review of individual NRSA fellowship awards (F30, F31 & F32), as well as the mentored career development awards (K08, K23 & K99/R00). Drs. Janet Cyr and Shiguang Yang will lead the presentation.

Workshop #2: Early Stage Investigators (ESI) and New Investigators (NI).

If you have plans to submit an R01 application for the first time, or have been unsuccessful with past R01 applications, then this workshop is for you. We will begin with a quick overview on how grant applications are processed including: assignment to NIDCD; preparation for review at different study sections; and when to contact staff during this process. Our goal is to provide information that facilitates a successful grant submission as a new (NI) or early stage investigator (ESI), use of the NIH Research Grant (R01), Exploratory/ Developmental Research Grant (R21), SBIR/STTR applications from a small business, and the NIDCD Small Grant Award (R03). Other issues will be addressed during the question/answer session. Drs. Roger Miller and Kausik Ray will lead the presentation.

Conclusion

These two sessions are intended for graduate students, postdoctoral fellows, new clinician investigators, and Early Stage Investigators (ESI) and New Investigators (NI).

Available funding mechanisms appropriate for new investigators will be discussed.
Protective Mechanism of the Outer Hair Cells from Traumatic Noise by Prior Whole-Body Heat Stress

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Background
The outer hair cell (OHC) motility results in cochlear amplification, leading to high sensitivity and sharp frequency selectivity of mammalian hearing. Unfortunately, however, OHCs are vulnerable to acoustic overstimulation. The OHCs have been reported to be protected from such traumatic exposure by prior sublethal conditioning, such as nontraumatic sound exposure, heat stress, etc. However, the mechanisms underlying conditioning-related cochlear protection remain unknown. In the present study, therefore, the effects of whole-body heat stress on the structure and function of OHCs were analyzed.

Methods
CBA/JNCrj strain male mice, aged 10–12 weeks (25–30 g), were used. The animals were anesthetized and placed in an aluminum boat floating in a hot water bath (46.5°C) to raise their rectal temperature up to 41.5°C. It was maintained at that temperature for 15 min. To evaluate the effects of heat stress on the structure of OHCs, Young’s modulus of OHCs, the amount of filamentous actin (F-actin) in OHCs and the expression level of heat shock protein 27 (HSP27) in the cochlea were measured before and after heat stress by atomic force microscopy (AFM), confocal laser scanning microscopy (CLSM) and Western blotting, respectively. To evaluate the effects of heat stress on the function of OHCs, distortion product otoacoustic emissions (DPOAEs) and the expression level of prestin in OHCs were evaluated before and after heat stress by an ER-10C acoustic system and CLSM, respectively.

Results
Heat stress caused an increase in Young’s modulus of OHCs at 3–6 h after its application (Fig. 1) along with an increase in the amount of F-actin (Fig. 2). Heat stress was also found to increase HSP27 in the cochlea and to enhance the DPOAE amplitude. On the other hand, the expression level of prestin was not changed by heat stress. These results suggest that heat stress induces HSP27 expression and thus increases F-actin in OHCs, increasing their stiffness, resulting in protection of the ear against noise induced hearing loss.

Conclusion
Conditioning with heat stress structurally modifies OHCs so that they become stiffer due to an increase in the amount of F-actin. As a consequence, OHCs possibly experience less strain when they are exposed to loud noise, resulting in protection of mammalian hearing from traumatic noise exposure. In contrast with F-actin, heat stress did not affect the amount of prestin.
Fig. 1. The mean and standard deviation of Young's modulus of the apical-turn OHCs in the control group (n = 10) and the anesthesia + heat groups with 3-h (n = 13), 6-h (n = 5), 12-h (n = 8), 24-h (n = 7) and 48-h (n = 12) intervals. Statistical analysis indicated significant differences between the control group and the anesthesia + heat groups with 3-h and 6-h intervals, as shown by asterisks (P < 0.05 by Student's t-test).

Fig. 2. The mean and standard deviation of the normalized average intensities of Fos-sin labeling in the control group (n = 13) and the anesthesia + heat groups with 5-h (n = 10), 6-h (n = 8), 12-h (n = 6), 24-h (n = 10), 48-h (n = 10) and 96-h (n = 8) intervals. Statistical analysis indicated significant differences between the control group and the anesthesia + heat groups with intervals of 3 h, 6 h, 12 h and 24 h, as shown by asterisks (P < 0.05 by Student's t-test).
Dose-dependent D-methionine Administration Significantly Reduces Permanent Impulse Noise-Induced Hearing Loss (NIHL) in Chinchillas

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Background
In previous animal studies, D-methionine (D-met) demonstrated otoprotection from drug-induced (cisplatin and aminoglycosides) and NIHL secondary to steady state noise exposures. In small scale Phase 1 and 2 clinical trials, D-met also protected from radiation-induced oral mucositis and cisplatin-induced hearing loss. D-met markedly reduced NIHL in animals when administered prior to and during steady state noise exposures or when first started 1 to 7 hours after these noise exposures. The purpose of this study is to determine the lowest D-met dose that is maximally effective in preventing impulse noise-induced hearing loss with noise exposures analogous to repeated M16 rifle fire. Currently we are preparing for clinical trials with the Department of Defense to prevent hearing loss for impulse noise exposures. If D-met prevents or reduces impulse noise-induced hearing loss, it could attenuate one of the most common causes of NIHL in soldiers.

Methods
Cohorts comprised five groups of 3 year old male Chinchillas laniger with ten animals per group. Groups received intraperitoneal injections of D-met at 0 (saline control), 25, 50, 100, or 200 mg/kg/dose every 12 hours beginning 2.5 days before and ending 2.5 days after noise exposure. Animals were exposed to simulated M-16 weapon fire at 155 dB peak SPL (150 repetitions at 2/s).

ABR thresholds were tested at baseline and on post-noise exposure day 21. Left and right ear thresholds were determined using tone-bursts centered at 2, 4, 6, 8, 14, and 20 kHz frequencies. Cochleae were harvested after the 21 day ABR and percentage of remaining outer hair cells measured.

Results
Significant reduction in ABR threshold shift at p≤ 0.05 occurred at 8 and 20 kHz for 100 and 200 mg/kg/dose groups and at p≤ 0.01 for 14 kHz in the 100 mg/kg/dose group. Hair cell analyses are in progress.

Conclusion
D-met can significantly reduce ABR threshold shift resulting from simulated M-16 weapon fire over a range of auditory frequencies and D-met dosing levels with 100mg/kg appearing to be optimal.

NIHL reduces troop redeployment and survivability /lethality on the battlefield. Current US military costs exceed 2 billion dollars annually D-met may be able to reduce NIHL in our military populations.

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Auditory Phenotypes of 129S6 and CBA Mice after High-Intensity Noise Exposures

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Background
Noise-induced hearing loss (NIHL) is a major health problem, and its effects are debilitating and irreversible. Using genetically homogenous mice as our animal model reduces inter-animal variability, and enables us to systematically assess inter-strain differences in vulnerabilities to NIHL. These phenotypic differences can be utilized to isolate the underlying genes involved. The inbred mouse strain 129S6/SvEvTac (S6) has been reported to be a remarkably noise resistant (NR) strain when exposed to noise levels that cause significant permanent threshold shifts (PTS) in the good hearing CBA/CaJ (CB) strain. Hirose and Liberman, 2003, assessed the noise exposure damage present when CB were exposed up to 116 dB SPL, which we use for comparison. Here we aimed to characterize the true extent of the NR phenotype in S6 by examining the effects of severe high-intensity exposure levels that cause dramatic PTS.

Methods
30 day old male unrestrained and unanesthetized S6 or CB mice were exposed to an octave-band noise (8-16 kHz) at high intensities (109 dB SPL, 115 dB SPL, or 118dB SPL) for two hours. A pre-conditioning stimulus was given immediately prior in order to avoid audiogenic seizure and death. Noise-induced PTS were assessed using distortion product otoacoustic emissions (DPOAEs) and auditory brainstem responses (ABRs). Mice were tested one or two days
prior to noise exposure for baseline, then again one and two weeks after the exposure. Inner ears were harvested after the final tests. Cochleas were then fixed and the organ of Corti was isolated from its bony confinement by fine whole-mount dissections. The tissue was then stained for immunohistochemical studies.

Results
After exposure to 118 dB SPL, the most intense acoustic trauma delivered in this study, both S6 and CB mice experienced 60-80 dB threshold shifts. Halving the power to 115 dB SPL resulted in a similar severe PTS in CB, but caused S6 mice to split into two different sensitivities. Half of the S6 pool endured a dramatic PTS, but the other half experienced partial acoustic protection. Hearing loss from 109 dB SPL exposure was less traumatic. The pattern of hair cell loss seen at the histological level is consistent with the degree of injury seen.

Conclusion
Our findings characterize the striking noise resistance phenotype of S6 mice by delivering noise exposures more intense than those used to date to study this strain.

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Mitochondrial peroxiredoxin 3 regulates hair cell survival in the cochlea
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Background
Mitochondrial peroxiredoxin 3 (Prx3) regulates intracellular redox balance and stress-induced apoptosis. Cochlear damage induced by traumatic noise, aminoglycosides and aging has been suggested to be linked to reactive oxygen species, disturbing cellular homeostasis. We therefore investigated the role of Prx3 in acquired hearing loss.

Methods
Auditory brainstem responses as measure of auditory function - Anti-Prx3 staining of surface preparations to determine expression of Prx3 in outer hair cells - Delivery of Prx3 siRNA to the round window via intra-tympanic application - Prx3 protein and mRNA levels from organ culture explants measured by Western blot and quantitative RT-PCR - hROS formation by fluorescence staining.

Results
In vivo, Prx3 increased transiently in mouse cochlear hair cells after traumatic noise, kanamycin treatment, or with progressing age before any cell loss occurred; when Prx3 declined, hair cell loss began. Maintenance of high Prx3 levels with the radical scavenger 2,3-dihydroxybenzoate prevented kanamycin-induced hair cell death. Conversely, reducing Prx3 levels with Prx3 siRNA increased the severity of noise-induced trauma. In explants of the murine organ of Corti, reactive oxygen species and levels of Prx3 mRNA and protein increased concomitantly at early times of gentamicin challenge. When Prx3 levels declined after prolonged treatment, hair cells began to die. The radical scavenger p-phenylenediamine maintained Prx3 levels and attenuated gentamicin-induced hair cell death. Gentamicin treatment also elevated levels of the transcription factor c-Myc, which targets the Prx3 gene, while a c-Myc inhibitor abolished the effect of gentamicin on both Prx3 mRNA and protein.

Conclusion
Our results suggest that Prx3 is up-regulated via c-Myc in response to oxidative stress and that maintenance of Prx3 levels in hair cells is a critical factor in their susceptibility to inner ear insults.

The research project described was supported by R01 grants DC009222 (to SHS) and DC-03685 (to JS) from the National Institute on Deafness and Other Communication Disorders, National Institutes of Health.
Prestin Regulation in Residual Outer Hair Cells after Noise-Induced Hearing Loss

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Background
The outer hair cell (OHC) motor protein prestin is necessary for electromotility. Previously, we found that OHC prestin expression was higher in Tecta transgenic mice with hearing loss. In the present study, we sought to determine the effects of noise-induced hearing loss on prestin expression.

Methods
We exposed cohorts of 5-6 week-old CBA mice to white noise (4-20 kHz at 100 dB SPL) for 4 hours.

Results
We assessed the cochlear epithelium by immunolabeling for prestin and myosin VIIa, and observed the expected pattern of OHC loss at the basal end of the cochlea. We then quantified prestin and found a significant increase in the fluorescence intensity from residual OHCs within the basal region of noise-damaged cochleae. We used whole cochlea preparations to measure prestin protein and found that the prestin/myosin VIIa ratio was elevated 1.7 fold at 7 days after noise exposure. We also measured mRNA expression changes in noise-damaged mice at 0.5, 3, 7 and 28 days after noise exposure. Noise exposure caused a significant increase in the prestin/myosin VIIa ratio. This effect was first detected at day 7 and persisted through day 28. ABR and DPOAE thresholds were elevated 30-40 dB immediately after noise. While most of the threshold shift recovered within 3 days, there was additional gradually recovery over the next month. At the 12 kHz cochlear location, where no OHCs were lost, the threshold and sharpness of masked tuning curves demonstrated full recovery by day 7. At the 32 kHz location, where partial OHC loss occurred, a near-complete recovery of tuning curve threshold and sharpness was found by day 7.

Conclusion
Taken together, we conclude that OHC prestin mRNA begins to become up-regulated >3 days after noise exposure and that prestin protein and cochlear amplifier function improve in an associated fashion with it. Prestin regulation may serve to partially compensate for OHC loss.
Renexin as a rescue regimen for noise induced hearing loss
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Background
Renexin (SK Chemicals Co Ltd, Korea) is a new combination drug of ginko biloba extract and cilostazol, which is known to have neuroprotective effect, anti-apoptotic effect and blood flow improvement effect. To evaluate the effect of Renexin on rescuing noise-induced hearing loss, we compared hearing and cochlear morphologies among different concentrations of Renexin-administered mice groups and control groups after noise exposure.

Methods
Mice were exposed to white noise at 110-dB sound-pressure level for 60 minutes at the age of 1 month. Different concentrations of Renexin (90mg/kg and 180mg/kg) were administered per oral for seven consecutive days after noise exposure. Auditory brainstem responses and distortion product otoacoustic emissions (DPOAE), cochlear pathology including efferent nerve endings were evaluated and compared among Renexin groups and control groups (noise and normal controls). Western blotting was performed to observe α-synuclein expression in the cochlea.

Results
Compared with mice of noise control, 90mg/kg and 180mg/kg Renexin groups showed less severe hearing loss from low to high frequencies at 7 days after noise exposure and their effects of rescuing hearing loss were dose-dependent manner. Various cochlear morphologic studies showed the far less severe outer hair cell damages in 180mg/kg Renexin-group compared to noise control group. Even 90mg/kg Renexin group showed the rescuing effect on outer hair cells after noise exposure. Efferent nerves under outer hair cells were more prominent in Renexin-groups compared to noise control group. Immunohistochemistry and western blot assay of α-synuclein also demonstrated a stronger expression of this synaptic vesicular protein in Renexin groups in dose dependent manner.

Conclusion
Renexin seemed to rescue hearing loss and cochlear damage after noise exposure in mice. Further studies to investigate the mechanism of Renexin on rescuing noise induced hearing loss will be necessary.

The auditory system functions in a circadian manner.
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Background
Circadian rhythmicity is essential for many bodily functions. A master clock is found in the hypothalamus, the suprachiasmatic nucleus (SCN), and is known to synchronize and coordinate rhythms to regulate various physiological functions including metabolism, inflammatory responses, oxidative reactions, feeding, sleep-wake patterns and the hypothalamic pituitary adrenal axis. Since the sensitivity of the auditory system can be modulated by the hypothalamic pituitary adrenal (HPA) axis and glucocorticoid receptors our working hypothesis is that the auditory system may function in a circadian manner.

Methods
Circadian oscillations of the PER2 protein were performed using tissues obtained from knock-in PERIOD2::LUCIFERASE (PER2::LUC) transgenic mice with a C57/B6 background. Ethical permission was granted by Stockholm’s Norra Djurförsöksetiska Nämnd. Cochlear explants were placed under photomultiplier tube detector assemblies mounted inside a light tight incubator at 36.5oC. Acoustic trauma was free field broadband noise at 6 - 12 kHz at intensity of 100 dB SPL for 1 h. Auditory sensitivity was assessed with auditory brainstem response (ABR) thresholds for the frequency of 6, 12, 16 and 24 kHz. The equipment for the ASR apparatus consisted of two startle chambers (SR-LAB; San Diego Instruments, San Diego, CA).

Results
Functional auditory responses were found to vary with the time of day. The amplitude of the acoustic startle response was higher and latencies shorter during day compared to night. Moreover, a greater loss of auditory function was found when noise trauma was delivered at night. The isolated peripheral auditory organ, the cochlea, was shown to have a self-sustained circadian rhythmicity of the expression of the clock gene protein product PERIOD2 that changed during development. Glucocorticoids were shown to synchronize the rhythm of the mouse cochlea-clock and to kick-start
oscillations in the central auditory structures, cochlear nucleus and inferior colliculus. The expression of the interleukin-6 and the neurotrophic factor BDNF in the cochlea was differentially altered after noise trauma applied during day versus night, which may be associated with the subsequent hearing loss.

**Conclusion**
The fact that the auditory system shows circadian rhythmicity is an important finding with clinical importance since noise-induced hearing disorders are an important public health issue and are exponentially increasing in the human population.

**Auditory responses in bats show strong resistance to noise impairment compared to gerbils**

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**Background**
Echolocating big brown bats are active at night and use their hearing as the dominant sense for orienting, for finding food, and for navigating through complex surroundings such as dense vegetation. Their near-total, moment-to-moment reliance on hearing in the 10-100 kHz range makes them potentially vulnerable to hearing impairment from intense sound exposures and from aging. Their own sonar broadcasts are ~120-130 dB SPL, but there are known mechanisms that reduce self-stimulation to about 75-75 dB SPL per sound. However, they also fly frequently in their roosts or in swarms, where they experience prolonged, virtually continuous exposure to ~100 dB SPL levels from other bats without evident impairment of echolocation. Further, bats have a very long lifespan of 20 years or more and yet they retain their sonar-guided abilities in the wild and in laboratory psychophysical tests.

**Methods**
To explore the implications of the bat’s acoustic lifestyle on protection of hearing, we compared big brown bats and gerbils for the impact of noise exposure on the magnitude of neural responses (cochlear microphonics, local field potentials, evoked potentials) to short, wideband frequency modulated (FM) stimuli.

**Results**
In experiments with broadband noise exposure equated for hearing range (20-100 kHz for bats, 3-30 kHz for gerbils), auditory responses to all FM sounds in gerbils decline in amplitude by up to 20 dB 1 minute after exposure and remain low for at least one hour. Recovery is complete after 10-12 hours. In bats, auditory responses to downsweeping FM sounds decline by 3-5 dB at 1 minute following exposure but have fully recovered by 30 minutes. Responses to upsweeping sounds are not affected.

**Conclusion**
The bat’s critical reliance on hearing for echolocation may have led to enhancement of its auditory protective mechanisms over the entire lifespan.
Lack of brain-derived neurotrophic factor in the cochlea but not in the brain hampers inner hair cell synapse physiology, but protects against noise induced afferent fiber loss

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Background
The precision of sound information transmitted to the brain depends on the transfer characteristics of the inner hair cell (IHC) ribbon synapse and its multiple contacting auditory fibers. We found that brain derived neurotrophic factor (BDNF), so far assumed to maintain spiral ganglia neuron survival, differentially influences IHC characteristics in the intact and injured cochlea.

Methods
We generated conditional knockout mice in combination with exocytosis measurements, calcium currents, auditory brainstem responses, ABR wave analysis, high resolution confocal microscopy.

Results
Using conditional knockout mice (BDNFPax2 KO) we found that resting membrane potentials, membrane capacitance and resting linear leak conductance of adult BDNFPax2 KO IHCs showed a normal maturation. Likewise, in BDNFPax2 KO membrane capacitance (∆Cm) as a function of inward calcium current (ICa) follows the linear relationship typical for normal adult IHCs. In contrast the maximal ∆Cm, but not the maximal size of the calcium current, was significantly reduced in high frequency but not low frequency cochlear turns correlated with a loss of IHC ribbons in these turns and a reduced activity of the auditory nerve (ABR wave I). Remarkably, a noise-induced loss of IHC ribbons, followed by reduced activity of the auditory nerve and reduced centrally generated wave II and III observed in control mice, was prevented in equally noise-exposed BDNFPax2 KO mice. This did not occur when BDNF was deleted in a mice model that lead to deletion of BDNF in central brain neurons.

Conclusion
Data describe a differential beneficial and harmful role of BDNF in the intact, respectively injured cochlea with dramatic impact on central sound processing that are discussed in a more widespread context of brain disorders.

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Investigating Long-Term Recreational Noise-Induced Hearing Loss in an Adult Population.

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Background
There is a great deal of concern about the risk of noise-induced hearing disturbances caused by listening to loud music. While there is clear evidence to support short term hearing disruption in some circumstances (Zhao et al. 2010), there is little, and mixed, evidence to support longer-term or permanent damage (Meyer-Bisch 1996, Tambs et al. 2004). A large scale study specifically investigating long-term recreational music exposure and any relationship between loud music exposure and hearing loss, tinnitus or hyperacusis is reported.

Methods
Participants were invited to complete two tasks using their home computer: First, an online English language speech-in-noise test. This was a modified digit triplet test presented at this conference last year (Vlaming et al. 2012). This test was optimised to detect high-frequency hearing loss, such as that caused by noise exposure; Second, a newly developed online long-term retrospective self-report questionnaire. The questionnaire contained questions on participants’ hearing, otological history, and most importantly asked for a life-history of their exposure to loud music. We anticipated recall of such a life history would be difficult. The use of a questionnaire tool called a “calendar instrument” has been shown to be useful in some interviewer-administered retrospective surveys (eg. Belli et al. 2007). As no studies were identified where a calendar instrument was used in an online, self-report context, we also compared a questionnaire using the instrument to a “standard” questionnaire in a set of two experiments (n=212). In the main study, commencing in October 2012, self-
reported music exposure is compared to both hearing test result and self-rated hearing questions, whilst controlling for other relevant factors.

**Results**
In the comparison of the questionnaire types, participants rated the questionnaires as being equivalent for difficulty, ease of use, and accuracy of recall (p<0.001). They rated the calendar instrument as being more than moderately helpful. No difference in data reliability was detected, but significantly more exposure was reported using the calendar (p<0.05), a finding taken to indicate better recall of exposures in other calendar instrument studies. Further analysis is continuing in 2012.

**Conclusion**
In addition to the above data, we will present results of the main screening experiment that will show the relationship between loud music exposure and hearing disturbances.

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**One Critical Subcellular Damage That Switches Noise-Induced Temporary to Permanent Threshold Shift**

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**Background**
Noise is the most common occupational and environmental hazard. Noise-induced hearing loss (NIHL) is the second most common form of sensorineural hearing deficit, after presbycusis. Low levels of noise exposure cause temporary threshold shift (TTS) only but not permanent threshold shift (PTS). Although promising approaches have been identified for reducing both TTS and PTS, currently there are no effective medications to prevent NIHL. Development of an efficacious treatment has been hampered by unknown cellular and molecular pathways involved in NIHL.

**Methods**
CBA/CaJ mice at 2 months old were exposed for 2 hours to 8-16 kHz octave band noise at 90 or 93 dB sound pressure level (SPL). There were also a no-noise control group and a group treated with zonisamide (120 mg/kg) just before the 93 dB noise exposure. TTS and PTS were measured one day and two weeks after noise exposure, respectively. Whole-mount fluorescent immunostaining of presynaptic ribbons by C-terminal binding protein 2 (CtBP2) and postsynaptic glutamate receptor 2/3 (GluR2/3) patches were analyzed by confocal microscopy and quantified by the imaging analysis software Velocity and our own custom program.

**Results**
Compared to the noise exposure at 90 dB causing only TTS, PTS was observed after exposure to 93 dB noise. The number of IHC ribbons and GluR2/3 patches was only slightly decreased after 90 dB noise exposure (0.78 and 0.59, respectively, averaged across 5, 10, 20, 28.3, and 40 kHz regions); while a significant synaptic loss was observed after 93 dB noise exposure (3.44 and 3.31, respectively, on average). Prophylactic administration of zonisamide reversed this synaptic loss after 93 dB noise exposure back to the loss similar to the 90 dB noise. Most importantly, no PTS was observed in the zonisamide-treated 93 dB noise group with reduced synaptic loss. Zonisamide saved about two ribbon synapses between IHC and SGNs on average across five frequencies examined.

**Conclusion**
Differences in cellular damage between TTS only and PTS have been observed, however, it was unknown what kind of cellular damage was critical for this difference. We have discovered that one critical difference between TTS only and PTS is a significant loss of synapses between inner hair cells (IHCs) and spiral ganglion neurons (SGNs) in cochleae with PTS, and this cellular damage can be prevented by a T-type calcium channel blocker (i.e., zonisamide).

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**Mmp7 knock-out potentiates noise-induced functional loss of the cochlea**

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**Background**
Metalloproteinases (MMPs), a family of zinc-dependent endopeptidases, play an important role in degrading the extracellular matrix. We recently reported the involvement of this gene family in noise-induced cochlear pathogenesis. The
study showed that Mmp7 was not constitutively expressed in the rat cochlear sensory epithelium, but was significantly upregulated after noise exposure. To investigate the role of Mmp7 in noise-induced cochlear pathogenesis, we examined the cochlear responses to acoustic trauma in Mmp7 knock-out mice and compared the response differences between the knock-out and the wild-type control mice.

Methods
B6.129-Mmp7tm1Lmm/J mice were used as experimental subjects and C57BL/6J mice were used as control subjects. Both the experimental and the control mice were exposed to a broadband noise at 120 dB SPL for 2 hours. Auditory brainstem responses were measured before and at various times after the noise exposure. After the final hearing test, the animals were sacrificed. The cochleae, kidney, liver and inferior colliculus were collected for transcriptional analysis of Mmp7 expression. The cochlear tissues collected were also used for pathological assessments.

Results
The expression of Mmp7 was not detected in any examined tissues (the cochlea, kidney, liver, and inferior colliculus) in the knock-out mice, but was detected in the kidney and liver in the wild-type mice, indicating that the knock-out mice lack Mmp7 expression. Under physiological conditions, the knock-out mice exhibited a similar threshold level of auditory brainstem responses as compared with the control animals, suggesting that Mmp7 knock-out does not affect normal hearing sensitivity. After the noise exposure, Mmp7 expression became detectable in the wild-type subjects, but remained undetectable in the knock-out mice, indicating that the Mmp7 knock-out mice lack the Mmp7 response to the acoustic overstimulation. As compared with the control mice, the knock-out showed a slightly greater hearing loss at 2 hours after the noise exposure (61 ± 19 dB for the wild-type mice vs. 67 ± 13 dB for the knock-out mice), but the difference was not statistically significant. Noticeably, the difference increased at 3 weeks after the noise exposure (37 ± 21 dB for the wild-type mice vs. 50 ± 12 dB for the knock-out mice, p < 0.01), suggesting that the lack of the Mmp7 response in the knock-out mice potentiates noise-induced hearing loss.

Conclusion
This study suggests that Mmp7 participates in the recovery process of noise-induced cochlear injury.

Isolation of the sensory cell-enriched, OHC-enriched and IHC-enriched samples from mouse cochleae with the high integrity of RNAs for transcriptional analyses
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Background
Analyses of molecular mechanisms of sensory cell degeneration rely on the collection of high quality of sensory cell samples from the cochlea. Traditional methods for collection of whole cochlear tissues or the sensory epithelium tissue are relatively easy, but the collected tissues contain a large quantity of non-sensory cells. The methods that are suitable for the collection of sensory cell-specific samples, such as in vitro hair cell isolation, a fluorescence activated cell sorting, and laser capture microdissection, are time consuming and unable to maintain the integrity of RNAs. The current investigation was designed to develop a microdissection technique for collection of high quality sensory cell-enriched samples from mouse cochlea for transcriptional analyses.

Methods
A microdissection technique was developed to isolate three types of cochlear samples from mouse cochleae: the sensory cell-enriched, OHC-enriched, and IHC-enriched samples. We also developed a method of image analysis for normalization of starting materials for transcriptional analyses. Using a pre-amplification technique, we examined the expression pattern of 12 reference genes (18SrRNA, Actb, Hprt, Hsp90ab1, Nono, Ppia, Rpl13a, Tbp, B2m, Ldhal6b, Gusb, and Tfrc) in the sensory cell-enriched samples, and identified the stable reference genes after exposure to a broadband of noise at 120 dB SPL for 1 hour.

Results
The reported method provided several advantages over the currently-available techniques. First, the cell types in each sample could be clearly defined. The sensory cell-enriched sample contained the hair cells and three types of supporting cells, Deiters cells, pillar cells and inner sulcus cells. The OHC-enriched sample contained OHCs, Deiters cells and pillar cells. The IHC-enriched sample contained IHCs, pillar cells and inner sulcus cells. Second, the integrity of RNAs extracted from the samples is well preserved. Third, the tissue could be collected from a defined cochlear site, suitable for the analysis of a site-specific change in gene expression. Using this technique for sample collection, we found expression of all 12 examined reference genes in the sensory cell-enriched samples. Among these genes, eight (18SrRNA, Actb, Hprt,
Hsp90ab1, Nono, Ppia, Rpl13a and Tbp) were relatively stable after exposure to the intense noise. Finally, we have successfully applied this method to sample collection from rat cochleae.

**Conclusion**
The reported microdissection technique is able to provide sensory cell-enriched samples with the well-preserved RNA integrity.
(Supported by NIH R01 DC010154)

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**Apoptosis and Molecular mechanism involved in cochlear implant trauma**

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**Background**
We described previously, the pattern of hearing loss post cochlear implantation (CI) and electrode insertion trauma (EIT) in animal models. This loss of hearing post CI has both an acute component (direct trauma) and an early post operative component (cellular and molecular damage) that develops over the period of at least a week following the initial trauma event. We demonstrated that insertion of a cochlear implant electrode array, not only causes direct tissue trauma and cell losses via necrosis, but also generates molecular events that will contribute further to a loss of residual hearing: e.g. oxidative stresses and release of pro-inflammatory cytokines that can lead to the initiation of programmed cell death; activation of the caspase pathway which can result in apoptosis; generation of pro-apoptotic signal cascades via, for example, mitogen-activated protein kinases/c-Jun-N-terminal kinases (MAPK/JNK); these may initiate programmed cell death within the damaged tissues of the cochlea. To gain a better understanding of the molecular mechanisms involved, we have investigated this process in a Guinea pig model of CI and tested the otoprotective effect of an inhibitor of JNK pathway.

**Methods**
Animals were exposed to EIT with and without treatment of JNK pathway inhibitor : AM-111 at round window. Immunostaining for HNE, activated Caspase-3, CellROX® and phospho-c-Jun were performed at 6hrs, 12hrs and 24hrs post-EIT. Neurofilament, Synapsin and FITC-Phalloidin staining for hair cells counts are performed at 90 days post-EIT. Guinea pig hearing thresholds were measured by ABR responses before and after cochlear implantation in four groups: EIT; pre-treated with hyaluronate gel ½ hr before EIT (EIT+Gel); pre-treated with hyaluronate gel/AM-111 ½ hr before EIT (EIT+AM-111), unoperated contralateral ears as controls.

**Results**
We observed an increase in ABR thresholds, post-EIT in the cochleae of EIT only and EIT+Gel treated animals. There was no significant increase in hearing thresholds in either EIT+AM-111 treated or unoperated control ears were observed. We observed protection of organ of Corti sensory elements (i.e. hair cells (HCs), supporting cells (SCs), nerve fibers and synapses) provided by AM-111 at 3 months post-EIT.

**Conclusion**
Molecular mechanisms involved in program cell death (PCD) of hair cells are different than the one involved in PCD of support cells. Local delivery of AM-111 prevented EIT-induced hearing, HCs losses and neural elements at 3 months.

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**Do Supernumerary Sensory Cells in the Cochleae of Rbl2 Null Mice Provide Protection from Noise-Induced Trauma?**

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**Background**
The Rbl2 null mouse is unique among mutant animal models characterized by the presence of supernumerary cochlear hair cells in that sensitivity to sound is normal in spite of the presence of an extra row of inner hair cells and extra rows of outer hair cells in the apical half of the cochlea (Rocha-Sanchez et al., 2011). The availability of a mouse model exhibiting extra sensory cells and normal thresholds creates an opportunity to determine if extra sensory cells offer protection from
hearing loss resulting from exposure to environmental noise that causes permanent hearing loss in control mice. The purpose of this study was to address this question directly.

Methods
A key element of the experimental design was to select potentially traumatizing noise bands that targeted cochlear regions populated with supernumerary sensory cells (the apical half of the cochlear spiral) and otherwise normal cochlear regions in the basal half of the end organ. Separate groups of Rbl2 null and wild-type (WT) mice were exposed to octave-wide bands of noise centered on 5.7 kHz or 11.3 kHz that were delivered for one hour at 120 and 110 dB SPL, respectively. Auditory brainstem responses (ABRs) to brief tone-bursts were used to estimate the magnitude of hearing loss produced by noise exposure, and to track recovery from hearing loss on a daily basis for the first four post-exposure days and at approximately 7, 14, 30 and 60 days thereafter.

Results
Immediately following exposure to traumatizing noise, ABR thresholds were elevated as much as 60 dB relative to pre-exposure, baseline values in both Rbl2 null and WT mice. Recovery of function progressed at approximately the same rate and was nearly complete for both Rbl2-deficient and control mice following exposure to the noise band centered on 11.3 kHz. However, Rbl2 null mice exposed to the noise band centered on 5.7 kHz recovered more completely than WT mice over the course of the first two weeks following exposure, but thresholds to tone-bursts increased in Rbl2−/− mice studied 30 and 60 days following noise exposure and threshold-frequency curve profiles observed at the two month post-noise exposure time point were similar to those observed in WT cohorts.

Conclusion
Findings reported here suggest that mice exhibiting supernumerary sensory cells are less vulnerable to consequences of noise trauma than WT cohorts in the short term, but not under long term conditions.

Methylene Blue Prevents Noise Induced Hearing Loss
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Background
The present study was performed to determine whether methylene blue, a powerful synthetic redox compound, can provide protection against noise induced hearing loss. Originally synthesized as a textile dye, methylene blue is currently being examined as a treatment for many human ailments including acute lung injury and Alzheimer’s disease. In low doses, methylene blue, which is highly cell permeable and readily crosses the blood-brain barrier, acts as an electron shuttle in mitochondria transferring electrons to oxygen thereby greatly reducing superoxide production. Additionally, methylene blue has been shown to directly inhibit the activity of both inducible and constitutive forms of nitric oxide synthetase (NOS).

Methods
Male CBA/CaJ (8 weeks old) mice (n=5/group) were injected with methylene blue (0.5 mg/kg IP) or water at 24 and 1 hr prior to exposure to the moderately damaging noise levels of 110 dB SPL, 4-45 kHz for 3 hrs. At 1 day, 1 week and 2 weeks, Auditory Brain Stem Response (ABR) was measured. The ABR to a 1-ms rise-time tone burst at 4, 8, 12, 16, 24, and 32 kHz was recorded and thresholds obtained for each ear. The effect of methylene blue on NOS and NADPH oxidase activities were measured using DAF-FM and CellROX Deep Red Reagent, respectively.

Results
The results revealed that prior treatment with methylene blue provided significant protection against noise induced hearing loss across all measured frequencies.

Conclusion
Here, we provide evidence that pre-treatment with low dose methylene blue can prevent hearing loss due to loud sound exposure. The mechanism behind this protection is likely the strong antioxidant properties of methylene blue that prevent loud sound induced free radicals from damaging the tissues of the inner ear. Supported by NIDCD DC 00105.
Auditory-based cognitive training improves the subcortical differentiation of stop consonants in older adults

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Background
Speech sounds are differentiated by acoustic cues which the auditory brainstem represents with tremendous fidelity. However, aging degrades the neural precision of subcortical speech encoding due to converging sensory and cognitive factors. Auditory training may prevent or reverse age-related declines in sensory processing. In animals, short-term perceptual training can reverse the effects of aging indexed by single neuron activity in auditory cortex.

Methods
We investigated the use of perceptual training to improve subcortical speech sound differentiation in older adults. We collected auditory brainstem responses to the stop consonants \([\text{ba}]\) and \([\text{ga}]\), which differ in their consonant-vowel (CV) transition formant trajectories, and so elicit brainstem responses which are out of phase in the transition period. After a first test, a group of older adults (ages 55-69) completed eight weeks of training which directed attention to speech transition cues by adaptively expanding and compressing CV transitions. After training we again collected brainstem responses.

Results
With training, subcortical consonant differentiation improved, indexed by an increase in the phase difference between responses in a time and frequency band corresponding to the CV transition. These changes in physiology were accompanied by improvements in speech perception, notably, hearing in noise. An active control group showed no changes.

Conclusion
In older adults, perceptual training can induce plasticity at fundamental stages of auditory processing. Specifically, training can improve the fidelity with which the brainstem transcribes rapidly-changing speech cues, which are especially vulnerable to the effects of aging. This plasticity can instill real-world communication benefits for listening in everyday situations. Supported by NIH (RO1 DC01510) and the Knowles Hearing Center.
Training reverses central processing effects of sensorineural hearing loss in older adults

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Background
Aging is accompanied by a loss of sensory function, and hearing impairment can be especially devastating due to reduced communication ability. Older adults report that speech, especially in noisy backgrounds, can be uncomfortably loud yet still unclear. Improving audibility through amplification provides only a partial remedy; impaired speech processing may limit the benefits of increased auditory input. Hearing loss may result in enhanced encoding of the slowly varying envelope (contributing to perceived loudness), dominating the response to the extent that the details of the rapidly varying fine structure (contributing to perceived clarity) are less salient. We hypothesized that older adults with hearing loss can be trained to reweigh envelope cues relative to the temporal fine structure.

Methods
We recruited older adults (ages 55 to 79) who were randomly assigned to an experimental or an active control group, both of which completed 8 weeks of computer-based auditory/cognitive training. All participants were tested pre- and post-training with electrophysiological, perceptual, and cognitive measures. We compared the effects of training in subgroups of participants with normal hearing and hearing impairment.

Results
After training, the experimental group with hearing loss had improved subcortical speech-in-noise processing. In particular, the relative representation of the envelope to the temporal fine structure was restored to that of individuals with normal hearing. No changes were noted in the active control group. Importantly, improvements in speech encoding were related to improvements in speech perception.

Conclusion
Central processing deficits associated with hearing loss can be restored through training. The speech-evoked brainstem response can be used as an index of these changes. This work is supported by the NIH (RO1 DC01510) and the Knowles Hearing Center.

Medial Olivocochlear Efferents: Changes in Shaker-2 Mice with Congenital Deafness

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Background
One of the main consequences of hearing loss is the impaired ability to understand speech in noisy environments. Medial olivocochlear (MOC) efferents are believed to be involved in this process due to their cholinergic damping effects on outer hair cell (OHC) function. Because deafness represents an extreme form of hearing loss, we examined changes in MOC neurons of hearing and deaf shaker-2 mice using immunohistochemistry for the enzyme Choline Acetyltransferase (ChAT). Differences in MOC cell features observed between these two groups of animals will guide future studies on milder forms of hearing loss.

Methods
ChAT staining was performed in hearing and deaf shaker-2 mice at various ages, and changes in immunopositive cell size and number were quantified. Additionally, some animals received injections of fluorogold into the round window of the cochlea to retrogradely label MOC neurons. Double labeling was used to determine what proportion of immunopositive cells of the superior olive were in fact MOC efferents.

Results
By 3 months of age, the OHCs of deaf shaker-2 mice have completely degenerated in the basal region of the cochlea (Gong et al., 2006). At this age, our data reveal that the MOC cells of deaf shaker-2 mice are significantly smaller (~20%) than those of age-matched hearing heterozygotes. By 10 months of age there is noticeable MOC cell loss and a ~50% reduction in cell size compared to normal hearing mice. These preliminary results indicate that MOC cell size and number progressively decreases with duration of deafness.

Conclusion
MOC efferents are implicated in the modulation of afferent input by regulating OHC activity, suggesting a critical role in extracting sound signals from noise. Abnormalities in MOC cell characteristics that accompany deafness indicate that
central auditory dysfunction is linked to peripheral damage. Furthermore, the pathology of MOC efferents is progressive, suggesting a use-dependent deterioration. It remains unclear, however, if this change is due to a loss of afferent input to MOC cells or the loss of their peripheral targets. Future studies will examine changes in MOC cells in non-congenital forms of hearing loss beginning at different ages in order to determine which factors most contribute to the degradation of this reflexive pathway.
Synaptic Transmission at Endbulb of Held Synapses During Age-Related Hearing Loss in CBA/Caj Mice
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Background
Age-related hearing loss (AHL) is a slow process in which peripheral hearing sensitivity diminishes with age. The changes in the central auditory system, particularly in the cochlear nucleus, however, are less clear. Here, we investigated synaptic transmission at the endbulb of Held synapse between the auditory nerve and cochlear nucleus neurons during AHL.

Methods
CBA/CaJ mice bred in–house were reared under standard vivarium conditions in microisolater cages until 31-38 days old, 85-105 days old, and 15-19 months old. Auditory brainstem evoked responses (ABR) to clicks were measured in each animal prior to preparation of brain slices. Voltage-clamp recordings were then made from bushy cells in parasagittal slices of the cochlear nucleus to measure the excitatory postsynaptic currents (EPSC) evoked by auditory nerve stump stimulation.

Results
The ABR showed ~10 dB SPL increase in hearing threshold in 15-19 month old mice relative to the two younger groups. However, the amplitude of the 1st ABR waveform increased more slowly with increasing intensity in both older groups, suggesting that hearing loss was present even in the 85-105 day old animals in spite of normal thresholds. When stimulating the auditory nerve in slices, we found that the EPSC amplitude in bushy cells was reduced while the paired-pulse ratio was increased in the oldest group. Furthermore, there was a significant increase in asynchronous release at the end of 400 Hz trains in the oldest group.

Conclusion
CBA mice exhibit decreased ABR amplitudes by 100 days of age, in spite of near-normal thresholds. Furthermore, at the first central site of synaptic transmission, AHL is associated with both weaker synaptic transmission (decreased release probability) and a compromised ability to sustain temporally precise transmitter release.

Unilateral Cochlear Implant Stimulation Maintains the Molecular Balance of the Plasticity Marker GAP-43 in Auditory Brainstem Circuits
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Background
In early life, millions of synapses develop in the mammalian brain. At this stage, high levels of phosphoprotein GAP-43 were detected in neuronal somata, axons and immature synapses. With maturation of the nervous system, synthesis of this protein decreased in most neurons, including those of the central auditory system. However, some regions maintain levels of GAP-43 mRNA, among them the lateral superior olive (LSO) and the central inferior colliculus (CIC). An additional region is the hippocampus, known for its involvement in learning. It is still unknown what happens molecularly in the brain if one ear or both fail in adulthood? Can a cochlear implant (CI) help to replace the missing input on the
molecular level? This study investigated the correlation between GAP-43 mRNA expression and symmetry of bilateral sensory inputs within the auditory system of rats.

Methods
For quantification, GAP-43 mRNA levels were determined in LSO and CIC. To create different auditory input conditions, we worked with four experimental groups of young adult, wild-type Wistar rats. (1) Rats with a unilateral CI, stimulated for up to 7 days, rats with unilateral (2) or bilateral (3) CI(s) for up to 10 weeks without stimulation, and (4) normal hearing controls.

Results
We found that GAP-43 mRNA levels shifted as a function of activity balance and activity patterns. The control group (4) and the unstimulated rats with bilateral CIs (3) had almost identical GAP-43 mRNA levels on both sides of the brain in LSO and CIC. By contrast, mRNA levels were significantly unbalanced for the unstimulated group with only one CI (2). In contralateral LSO, GAP-43 mRNA expression increased as compared to control level. Ipsilateral CIC reached significance against control level after long-term implantation of 3 or 10 weeks. By monaural CI stimulation (1), GAP-43 mRNA levels were left-right symmetrical for LSO and CIC. Still, levels were higher than in control rats in LSO for all stimulation times and in CIC by 7 days of EIS.

Conclusion
Overall, our data suggest that GAP-43 is more than a marker for neurite outgrowth and early stages of synaptogenesis. When activity input is different between left and right ear of adult rats, this imbalance is displayed by levels of GAP-43 mRNA expression. Remarkably, the balance can be completely sustained if the hearing impaired ear became activated by simple stimulation patterns by way of a CI.
Presynaptic ion channels regulate quantal size in the calyx of Held

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Background

Synaptic vesicles accumulate transmitter using energy from the proton electrochemical potential ($\Delta \mu_{H^+}$), composed of an electrical potential ($\Delta \psi$) and a chemical gradient ($\Delta \text{pH}$). Previous studies suggested that glutamate uptake into synaptic vesicles by the vesicular glutamate transporter (VGLUT) is dependent mostly on $\Delta \psi$ rather than $\Delta \text{pH}$. Recently, a study showed that rat brain synaptic vesicles express a K$^+$(Na$^+$)/H$^+$ monovalent cation exchanger activity that converts $\Delta \text{pH}$ into $\Delta \psi$ and promotes synaptic vesicle filling with glutamate (Goh et al, 2011). Manipulating presynaptic K$^+$ at the calyx of Held, a giant glutamatergic terminal in the auditory brainstem, influenced quantal size, indicating that synaptic vesicle K$^+$/H$^+$ exchange regulates glutamate release and synaptic transmission. Although the intracellular K$^+$ concentration is relatively stable, the Na$^+$ concentration may fluctuate widely during spike activity, raising the question of whether Na$^+$ may have a regulatory influence on VGLUT activity.

Methods

Whole-cell electrophysiology and Na$^+$-imaging recordings were made from the calyx of Held in brainstem slices of P8-14 rats.

Results

Our experiments showed that intracellular Na$^+$ strongly affects glutamate uptake: increasing Na$^+$ in the presynaptic recording pipette potentiated mEPSC amplitude (quantal size) while decreasing Na$^+$ in the pipette reduced mEPSC. We found that hyperpolarization-activated cyclic nucleotide-gated (HCN) ion channels control the resting Na$^+$ concentration and that blocking HCN channels with Cs$^+$ reduced the amplitude of mEPSCs. Moreover, modulating HCN channels with forskolin or 8-Br-cAMP increased mEPSC size.

Conclusion

Plasma membrane ion channels control intracellular Na$^+$ and therefore regulate glutamate uptake and synaptic transmission. Because high-frequency spike activity leads to Na$^+$ accumulation in terminals, we expect activity dependent facilitation of glutamate uptake will speed vesicle replenishment and sustain synaptic transmission.
Synaptic and Membrane Properties of Neurons in the Ventral Nucleus of the Lateral Lemniscus (VNLL) of Mice

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Background
Neurons in the lateral lemniscus receive numerous inputs from lower auditory brainstem nuclei and send prominent inhibitory projections to the inferior colliculus. Based on the anatomical inputs and physiological properties, neurons in the lateral lemniscus can be subdivided into at least two distinct nuclei, the dorsal (DNLL) and the ventral nucleus of the lateral lemniscus (VNLL). Whereas properties of DNLL neurons have been well described, less is known about neurons in the VNLL. Neurons in the VNLL receive excitatory inputs from the contralateral ventral cochlear nucleus (VCN) and inhibitory inputs from the ipsilateral medial nucleus of the trapezoid body (MNTB). However, little is known about the physiological properties of these neurons. Therefore, we started to characterize the membrane and synaptic properties of neurons in the VNLL of mice.

Methods
Biophysical membrane properties and synaptic inputs were characterized by whole-cell patch-clamp recordings of VNLL neurons in acute brain slices from P10/11 and P17/18 C57Bl6J mice. Excitatory and inhibitory synaptic currents were evoked by stimulating the incoming fibers of the lateral lemniscus with a glass electrode placed ~50 µm ventral to the recorded neuron. Underlying synaptic receptors were determined by pharmacological blockade of the respective receptors. VNLL neurons were identified by their dorsal/ventral position within the lemniscal fiber tract. Only neurons in the most ventral part in the fiber tract were analyzed.

Results
VNLL neurons predominantly displayed an onset-type firing pattern in response to depolarizing current step injections (1s). During hyperpolarizing current injections these neurons exhibited a prominent depolarizing voltage sag. This indicates that both low threshold K⁺-channels and HCN-channels are expressed in the VNLL.

Stimulating the incoming fibers evoked both excitatory and inhibitory synaptic responses. In P17/18 mice excitatory synaptic currents were mediated by AMPA and NMDA receptor. Both AMPA and NMDA receptor currents had large amplitudes. Inhibitory synaptic responses had fast time constants (around 2 ms) and were largely mediated by glycine at postnatal day 17/18.

Conclusion
Our results show that synaptic and membrane properties of VNLL neurons, that are located in the lemniscal fiber tract, are fairly uniform. These neurons receive large excitatory and inhibitory inputs and display membrane properties that ensure precise integration of excitatory and inhibitory inputs.

Competing mechanisms of short-term synaptic plasticity in the chick cochlear nucleus

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Background
In the auditory system, stimuli of different frequencies are processed in parallel frequency-tuned circuits, beginning in the cochlea. Activity of avian auditory nerve fibers reflect this frequency-specific topographic pattern, known as tonotopy, and impart frequency tuning onto their postsynaptic target neurons in Nucleus Magnocellularis (NM). While all NM neurons perform the basic function of encoding the timing properties of acoustic stimuli, physiological specializations exist along the tonotopy that reflect the characteristic frequency (CF) of their auditory nerve fiber inputs. One known feature of synaptic physiology that has not been investigated across the tonotopic axis of NM is short-term synaptic plasticity.

While the utility of comparing synaptic plasticity across different neurons in different brain regions is limited, NM offers a rather homogenous population of neurons with a distinct topographical distribution of synaptic specialization that is ideal for the investigation of short-term synaptic plasticity. We have previously demonstrated that synapses with a high CF are more resistant to short-term synaptic depression, which is due in part to a larger readily releasable pool (RRP) and a faster rate of recovery than their low CF counterparts. In this study, we examined the contribution of current amplitude and postsynaptic mechanisms to the observed difference in depression.

Methods
We used in vitro whole-cell patch clamp with chickens aged E19-P1. Brainstem slices were collected, and slice order determined tonotopic position of NM neurons. A stimulating electrode was placed in a position proximal to the cell until a
unitary evoked excitatory postsynaptic current (EPSC) was observed. Cyclothiazide (40 μM) was added to aCSF to evaluate receptor desensitization.

Results
Superimposed on the tonotopic distribution of synaptic depression, we show that larger inputs show more depression, regardless of characteristic frequency. Experiments with cyclothiazide showed that larger EPSCs also caused larger AMPA receptor desensitization. However, this postsynaptic effect was small relative to presynaptic components of short-term synaptic depression, and was not sufficient to explain the tonotopic distribution of depression.

Conclusion
Across the tonotopy of NM, high CF synapses are more resistant to depression than their low CF counterparts. We show that larger EPSCs exhibit greater depression, even within frequency regions. In addition, the magnitude and distribution of postsynaptic receptor desensitization did not reflect the difference in short-term synaptic depression across the tonotopy. Therefore, the presynaptic mechanisms of RRP size and vesicle recovery are the most likely mechanisms that underlie synaptic specialization in NM.

How does Jaundice Cause Deafness in a Model Mouse?
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Background
Acute bilirubin encephalopathy (ABE) and Chronic bilirubin encephalopathy (or kernicterus) are serious neurological events with an incidence of 1/100,000 and 1/150,000 in newborns in the United Kingdom. Both conditions are associated with hyperbilirubinemia. Jaundice- (Maisels, Early Human Development, vol. 85 pp 727-732, 2009) where increased plasma levels of free bilirubin cross the blood-brain-barrier and cause neuronal damage in the CNS (Haustein et al, J. Physiol., Vol. 588.23 pp 4683-4693, 2010). The link with vulnerability of the auditory system is well documented, but the mechanism(s) of bilirubin neurotoxicity remain unclear. To investigate damage in the brainstem auditory pathway we focused on a giant synapse that has a pivotal role in the circuitry involved in sound localization. This synapse, between the anterior ventral cochlear nucleus (aVCN) and the neurons of the medial nucleus of the trapezoid body (MNTB), is known as calyx of Held (Kopp-Scheinplug et al, Hearing Res., vol. 279 pp22-31, 2011)

Methods
Hyperbilirubinemia was induced in CBA mice by a single intraperitoneal injection of free bilirubin (0.5 mg/g and Sulfadimethoxine 0.3mg/g) and the auditory function was monitored by recording in vivo auditory brainstem responses (ABR) 4h-24h and 5 days afterwards. We used an in vitro slice preparation to study transmission at the calyx of Held combined with multi-photon imaging and immunohistochemistry for the presynaptic marker VGLUT2 (vesicular glutamate transporter 2). Additionally the acute effect of bilirubin was investigated on slices from untreated animals, using patch clamp recording.

Results
We showed a deficit of ABRs at 4h which had recovered by 24h and 5 days after injection. Using multi-photon imaging and immunohistochemistry for VGLUT2 we found disruption of the synapses at the calyx of Held at both 4h and 24h after injection. Patch clamp recordings showed a decrease in the threshold for the firing of action potentials in the MNTB, after acute perfusion of the slices with free bilirubin. In control experiments, bilirubin caused no change in the amplitude of evoked EPSCs mediated by AMPA or NMDA receptors.

Conclusion
We conclude that this mouse model for hyperbilirubinemia exhibits an acute damage to the auditory system, involving a disruption of the synapses at the level of calyx of Held and provides a good model for further understanding mechanisms of injury caused by jaundice.

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Intrinsic firing heterogeneity in the cochlear nucleus improves population coding of temporal fluctuations during noisy current stimulation

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Background

There is widespread evidence for intrinsic property diversity in many brain areas, but the role of such diversity in neural coding is not well understood. While neurons in the timing pathway of the avian auditory brain stem are fairly stereotyped in their firing properties, those in the intensity pathway show a broader range of biophysical diversity. Previous research investigated how the intrinsic properties of the cochlear nucleus angularis (NA) influenced the response to noisy current injections during recordings from slices in vitro that simulated the arrival of many nerve inputs. All physiological subtypes of NA neurons showed reliable, stimulus-locked firing to fluctuating noisy currents, but the degree of spike train reliability and precision correlated with cell type (Kreeger et al., 2012, J Neurophysiol.) In this study, we investigated how well the stimulus was encoded by individual NA neurons, how the diversity of intrinsic properties affected encoding and whether this diversity led to better population coding of the stimulus.

Methods

Patch-clamp recordings were made from chick (E17-18) brain stem slices while delivering filtered Gaussian white noise current stimuli. We used reverse correlation methods to reconstruct an estimate of the stimulus using the spike-triggered averages (STA) and assessed by cross-correlation with the original stimulus (low-pass, 300 Hz).

Results

To compare reconstructions across neurons with different properties, identical noise stimuli were presented during separate recordings from 60 NA neurons. Reconstruction estimates showed high degree of correlation with the original stimulus (0.50±0.08). STAs generated by two, uncorrelated noise stimuli were very similar within a given neuron, but differed across neurons for the same stimulus, suggesting that each neuron had a unique sensitivity to the current fluctuations. Comparing spike trains across neurons to identical stimuli, differences were observed in number and timing of spikes, but there were also many correlated, precise spike events. Combining information from spike trains drawn from the population improved stimulus reconstruction estimates (0.67±0.02, 40 cells).

Conclusion

Temporal fluctuations in current inputs were efficiently coded by individual NA neurons and combining information across the population improved coding due to biophysical diversity. Highly reliable firing has also been observed in NA in vivo during repeated broadband noise stimuli (Steinberg and Pena, 2011 J. Neurosci. v31:3234), suggesting temporal envelope fluctuations may be encoded via the intensity pathway. Whether intrinsic property diversity contributes to envelope coding in vivo remains to be seen.

Development of Kcna1 Dominant-negative Lentivectors in the Central Auditory System

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Background

Throughout the nervous system, potassium channels are required for efficient signaling and action potential timing. In the central auditory system, the voltage-activated, delayed rectifier Kv1.1 subunit, encoded by the Kcna1 gene, is important to sustain the firing rate and timing of action potentials via its regulation of a low-voltage activated current (IKL) (Brew et al 2003; Gittelman and Tempel 2006; Kopp-Scheinplug et al 2003). To study how altered Kv1.1 channels affect functioning in the central auditory system it is necessary to establish an animal model that has mutated channels in the auditory system only. As Kv1.1 is expressed throughout the central nervous system, a necessary means to achieve this specificity in the auditory system is via localized stereotaxic injections of lentiviral vectors. The main goal of this study is to generate and establish lentivectors with a dominant negative Kcna1 mutation as a means of studying desynchronized activity in the central auditory system.

Methods

Recombinant lentiviruses were developed to provide a range of interference with endogenous Kv1 subunits. GFP-expressing lentivectors were produced by introducing AU1-tagged Kv1.1 cDNAs encoding wildtype and mutant to the lentivector. For in vivo expression, lentiviruses were injected into the anteroventral cochlear nucleus (AVCN) of adult mice,
which projects monosynaptically to the medial nucleus of the trapezoid body (MNTB). AVCN location was confirmed stereotaxically and also by detection of multi-unit activity. GFP intensity and the density of transfected neurons in both the AVCN and MNTB were determined using immunocytochemistry.

Results
We have produced repeatable Kv1.1 dominant negative lentiviral injections in the AVCN of adult mice, and this resulted in transfection of both AVCN neurons and MNTB pre-synaptic calyces. Additionally, we established a timeline of integration of the mutant Kv1.1 subunit into the genome of AVCN bushy cells. Using antibodies to the AU1-tag, we determined the time required for mutant Kv1.1 subunits to integrate into the MNTB calyces of Held.

Conclusion
These results suggest that it is possible to use a Kcna1 dominant negative lentivector as a means to disrupt Kv1.1 channel activity in the central auditory system. After optimizing the lentivector injections, we can record single unit activity from MNTB principal cells to determine what effect disrupted Kv1.1 activity will have on the physiological responses of these cells. Support: NIDCD R01-DC02739 (BLT) and 5T32DC000018-30 (JLT).

Imaging synaptic zinc release in the dorsal cochlear nucleus with newly developed zinc sensors and chelators
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Background
Vesicular zinc exists at many glutamatergic synapses in the brain. This transition metal is loaded into presynaptic glutamatergic vesicles and is co-released with glutamate during synaptic transmission. The molecular layer of the dorsal cochlear nucleus (DCN) contains high levels of vesicular zinc. Our group has recently reported that synaptically-released zinc initiates endocannabinoid synthesis that subsequently modulates synaptic strength. However, the amount of synaptic Zn²⁺ that reaches the postsynaptic membrane during synaptic stimulation remains unknown, mainly due to lack of appropriate zinc sensors and chelators.

Methods
Here, we have used new generation of fluorescent zinc sensors and chelators to examine how much, and under what conditions zinc is released in the DCN.

Results
We have found ZnT3-mediated zinc release following electrical stimulation with zinc-sensitive fluorescent probes.

Conclusion
We provide evidence for ZnT3-mediated zinc release in the DCN.
Glutamate-activated currents in unipolar brush cells

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Background
Unipolar brush cells (UBCs) of the dorsal cochlear nucleus (DCN) and vestibular cerebellum receive glutamatergic mossy fiber input on a complex brush-like dendrite. Studies of cerebellar UBCs identified different subtypes of UBC based on immunohistochemical markers and physiological profiles. We have examined auditory UBCs, and compared them to cerebellar UBCs with respect to the physiology of the mossy input and glutamate sensitivity.

Methods
Patch-clamp recordings were made on coronal slices from DCN or parasagittal slices from vestibular cerebellum in P16-23 mice. A monopolar or bipolar glass electrode was used for extracellular stimulation. Glutamate was pressure ejected from a pipette positioned near the dendritic brush. Mouse lines included mGluR2-GFP and wildtype C57/Bl6. GABA_A and glycine receptors were blocked to prevent inhibitory transmission.

Results
EPSCs evoked by extracellular stimulation evoked NBQX-sensitive EPSCs with a profile similar to that reported by Rossi et al. 1995: a small, fast inward current followed by a very slow delayed inward current. Occasionally, a train of stimuli would result in a delayed outward current consistent with a glutamate-activated K⁺ current. However synaptic responses were painfully rare. To explore glutamate sensitivity of UBCs, we therefore applied brief puffs of 1 mM glutamate. A variety of response profiles were elicited in different cells, suggestive of different cell types. Overall the results indicated the presence of at least 3 glutamate receptor subtypes in UBCs. 1) AMPA receptors (defined by sensitivity to NBQX), which generated a small, rapid inward current followed by an outward current and lastly a slow inward current. In some cases the AMPA current was negligible, but in all cases the current was markedly increased by cyclothiazide, a blocker of AMPAR desensitization. 2) A slow outward current blocked by mGluR2 antagonist, consistent with a GIRK channel. 3) A slow inward current blocked by antagonists to mGluR1 receptors.

Conclusion
UBCs express an array of glutamate receptors which generate complex response profiles. The high expression of metabotropic glutamate receptors coupled to K⁺ channels and other cation channels could grant these cells with a unique role in processing multisensory signals in the structures where they are present.

Dependence on otoferlin (and background strain) of magnitude and convergence of auditory nerve inputs on cochlear nuclear bushy cells.

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Background
Even before the onset of hearing, patterned spontaneous activity that starts in the cochlea ascends through auditory pathway (Tritsch et al. 2007; Tritsch and Bergles 2010). Otoferlin acts as a calcium sensor at synapses of inner hair cells, mediating synaptic transmission. In otoferlin -/- mice, inner hair cells release little neurotransmitter throughout the life of the animal (Roux et al. 2006; Longo-Guess et al. 2007). Bushy cells in the anteroventral cochlear nucleus receive most excitatory input through large terminals, the end bulbs of Held. End bulbs in otoferlin mutants are smaller and more wispy (Wright et al., 2012). We compare the number of inputs and the magnitude of evoked excitatory postsynaptic currents (EPSCs) in bushy cells of otoferlin -/- mice with +/- and +/+ hearing control mice to learn how activity early in development affects end bulbs of Held.

Methods
Stimulation of bundles of auditory nerve fibers evoked EPSCs in bushy cells in slices that grew in steps of widely varying sizes with shock strenght, reflecting the recruitment of converging fibers (Cao et al., 2007; Cao and Oertel, 2010).

Results
The average number of steps was not different in deaf otoferlin -/- mice (4.2 ± 1.5 steps, n=36) and hearing +/- and +/+ controls (3.4 ± 1.9 steps, n=36) (p = 0.06) (P15-P22) suggesting that the number of inputs is not strongly affected by spontaneous and acoustically driven activity. Evoked EPSCs were enormous in all three genotypes, with maximal currents exceeding 60 nA in some cells. Unitary EPSCs in hearing +/- and +/+ mice were significantly smaller (5.2 ± 6.2 nA, n=36) than in deaf otoferlin -/- mice (7.6 ± 8.8 nA, n=36, p = 0.01). Because measurements of such large synaptic
currents are sensitive to series resistance compensation in patch clamp recordings, the measurements of amplitude must be viewed as approximate.

The unexpectedly large EPSCs in the +/+ C57BL/6J and C3HeB/FeJ background (maximal EPSCs 19.9 ± 12.2 nA, n=18), prompted a comparison of maximal EPSCs in various wild type strains recorded in our lab. We find large differences in the magnitude of maximal EPSCs between strains. In ICR mice maximal EPSCs were 2.7 ± 1.4 nA (n=23); in C57Bl/6 and 129S hybrids, maximal EPSCs were 1.0 ± 0.1 (n=3); in another hybrid strain, AKR, C57Bl6, LG, and BALBc, maximal EPSCs were 3.6 ± 1.4 nA (n = 6).

Conclusion
Activity and background strain matters.

Munc13 proteins differentially regulate short-term plasticity at the calyx of Held synapse in the mammalian auditory brainstem
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Background
Numerous molecular interactions at the active zone of synapses regulate the reliability of synapses to encode information over a wide range of frequencies in response to action potentials. In the CNS, the Munc13 gene family encodes critical molecules in the regulation of synaptic transmission and plasticity and consists of three genes, Munc13-1, Munc13-2 and Munc13-3, that express four Munc13 isoforms. Currently, it is unknown why despite the presence of Munc13-2 or Munc13-3, they are functionally dispensable at some synapses or why their loss in other synapses leads to increases in frequency-dependent facilitation. We addressed these questions at the calyx of Held synapse, a giant synapse in the auditory brainstem, that places a premium on speed and reliability of information transfer for auditory signal processing and allows for biophysical dissection of the different regulatory steps in CNS synaptic transmission.

Methods
Acute brainstem slices containing the MNTB were prepared from Munc13-2 KO, Munc13-3 KO, or Munc13-2-3 DKO or wild-type mice. Electrophysiology experiments to study AP-evoked EPSC’s were performed by afferent fiber stimulation in conjunction with patch clamp recording of the MNTB. To quantitatively understand the dynamics of the readily releasable pool of vesicles we performed paired recordings of the calyx/MNTB synapse. Quantitative immunohistochemistry experiments measuring Munc13 isoform levels at the calyx were done using the Munc13 XFP knockin animals. Experiments used mice before the onset of hearing (p9-11) or when the calyx is more functionally mature (p18-p21).

Results
We report that the calyx of Held synapse expresses 3 isoforms of Munc13 (Munc13-1, ubMunc13-2, and Munc13-3). Using detailed electrophysiological analyses of the Munc13 KO mice lines, we report that loss of both Munc13-2 and Munc13-3 leads to an increase in the rate of calcium-dependent recovery and a reduction in the readily releasable pool, but with no change in the fast releasing vesicle pool size but a selective loss of the slow releasing vesicle pool size. Using Munc13XFP knock-in mice, quantitative immunohistochemistry revealed that Munc13-1 is the dominant isoform. We also find that only the Munc13-1 isoform is highly colocalized with bassoon at the active zone.

Conclusion
Based on our data, we conclude that Munc13-2 and Munc13-3 isoforms compete with Munc13-1 to differentially regulate calcium-dependent recovery and the slow pool to fast pool conversion in central synapses.

Physiology of Golgi cell–granule cell synapses in the cochlear nucleus
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Background
The cochlear nucleus contains diverse granule cell domains conveying multimodal signals from mossy fibers to the molecular layer of the dorsal cochlear nucleus (DCN). Little is known about inhibitory cells that modify these multimodal signals at the level of the granule cell. A previous study (Balakrishnan and Trussell 2008) showed prominent GABAergic
and glycinergic inputs to granule cells in rat, but did not identify the sources of these inputs. We have used mouse lines in which GFP is expressed in inhibitory neurons and tdTomato is expressed in granule cells to facilitate identification of these cell types in different regions of the cochlear nucleus. Moreover, we explored modulation of inhibition by metabotropic receptors.

**Methods**

Single or paired patch-clamp recordings were made on slices from P16-23 mice. A monopolar or bipolar glass electrode was used for extracellular stimulation. Mouse lines included mGluR2-GFP, GlyT2-GFP, and GABA_A6-tdTomato. AMPA and NMDA receptors were blocked to prevent glutamatergic transmission.

**Results**

Extracellular stimuli reliably elicited IPSCs in granule cells from DCN and granule cell lamina. As in rat, IPSCs were mediated by GABA and glycine receptors, suggesting that presynaptic fibers released both transmitters. Paired recordings made from either GFP-expressing, non-unipolar brush cell neurons in mGluR2-GFP or GlyT2-GFP positive neurons synaptically coupled to granule cells revealed that at least one source of inhibition to auditory granule cells is local interneurons. For convenience, we collectively term these interneurons Golgi cells, although it is possible they represent a diverse class. Paired recordings always showed smaller IPSCs than with extracellular stimulation, implying that 3-4 Golgi cells converge on single granule cells.

A previous study demonstrated muscarinic and metabotropic glutamate receptor 2-mediated hyperpolarization of mouse Golgi cells (Irie et al., 2006). We confirm this report but also observed potent inhibition of evoked IPSCs in granule cells by carbachol or by LY354740. The effect of carbachol was mimicked by oxotremorine M, indicating the presence of muscarinic receptors. In paired recordings in which the presence of a presynaptic spike could be verified, synaptic inhibition persisted, indicating that inhibition is not generated by prevention of presynaptic spikes. Nevertheless, modulation of short-term plasticity of IPSCs by carbachol suggested that carbachol acted at a presynaptic site.

**Conclusion**

Golgi cells may control the efficacy of multimodal inputs to the DCN. Modulation of inhibition by metabotropic receptor systems may provide a means for state-dependent gating of these signals.

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**Noise-Induced Changes of Neuronal Spontaneous Firing Rates in Mice Cochlear Nucleus and Inferior Colliculus Brain Slices**

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**Background**

A noise trauma leads beside damage in the periphery to profound changes in the central auditory system. Recent work demonstrated that noise exposure has a strong and rapid impact on central neuroanatomy and neurophysiology. Despite an increase in apoptotic cell death and a subsequent reduction in cell densities, a single noise trauma is often followed by an elevation of spontaneous neural activity in auditory brain structures as demonstrated by electrophysiological recordings or activity-dependent imaging techniques. As most of these studies where performed in vivo, it remains unclear to what extend particular brain areas are responsible for the observed hyperactivity. Acute in vitro preparations provide the opportunity to investigate intrinsic properties of brain structures disconnected from ascending or descending neuronal input. It was the aim of the present study to investigate spontaneous firing activity at different time points after noise exposure in brain slices of the lower auditory pathway, namely the cochlear nucleus (CN) and inferior colliculus (IC).

**Methods**

Normal hearing mice (NMRI strain) were exposed to a broadband noise (5-20 kHz, 115 dB) for 3 hours under anesthesia and have been investigated immediately (acute group) or 7 or 14 days after trauma. Hearing thresholds were determined by auditory brainstem response audiometry. To investigate changes in spontaneous neural activity in vitro, electrophysiological single-unit recordings were carried out in the dorsal and ventral cochlear nucleus (DCN and VCN) and IC brain slices. Unexposed mice served as normal hearing controls.

**Results**

The ABR results demonstrate that hearing loss in the animals was highest immediately after noise exposure and recovered significantly within one week. However, a significant permanent threshold shift was present even at day 14 after noise exposure. Spontaneous neuronal activity was significantly increased in the acute group on the level of the DCN and VCN, no significant changes occurred in the IC. One week after noise trauma, firing rates did not differ from control
animals in any investigated structure. However, after 14 days, spontaneous activity was significantly elevated in the DCN as well as in the IC.

Conclusion
The data underline our previous findings that a noise trauma is acutely affecting central cellular properties and induces short-term plasticity, particularly in brainstem structures. Further, long-term physiological changes seem to appear slowly and might thus differ from the early effects, whereby the DCN and the IC (but maybe not VCN) are supposed to serve as key structures during this development.

Filter widths throughout the auditory system
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Background
Frequency selectivity is a fundamental and extensively studied property of the auditory system, yet it is not clear whether it is established at the periphery or if central mechanisms make a contribution. This is, in part, because two distinct approaches are taken, psychophysics and physiology, the results of which are difficult to compare. Psychophysical estimates are made using masking paradigms, whilst many physiological measurements in the past were made using pure-tone responses. Also, psychophysics measures detection thresholds, which are not necessarily comparable to the mean spike rates used in physiology.

Methods
To try and reconcile the differences between these two fundamentally different approaches, we measured auditory filter widths of single neurons, as well as multi-unit activity, in two brain regions; the inferior colliculus (IC) and primary auditory cortex (A1). This was performed in the anesthetized guinea-pig. Unlike most similar physiological experiments we used notched-noise masking of a pure-tone signal, which is more typically seen in psychophysics. Signal detection theory methods were used to convert spike firing rates into neurometric functions, similar to psychometric functions. Threshold signal to noise ratio values were calculated and roex(p, r) functions used to estimate filter size. In addition roex(p_l, p_u, r) functions were fit to pure-tone response areas, allowing a direct comparison to the masking approach.

Results
In the IC notched-noise results were relatively homogeneous and corresponded well with previous experiments in the literature, which used more traditional pure-tone methods and in more peripheral auditory brain regions. Simultaneously masked bandwidth estimates were significantly broader than forward masked ones, which concurs with results in psychophysics. However, pure-tone derived estimates were significantly broader than notched-noise ones calculated within the same unit (by a factor of ~2). In A1 pure-tone calculated bandwidths were equally as broad, but more surprisingly, notched-noise bandwidths were narrower than seen in the IC and behaviour.

Conclusion
Frequency selectivity seems to be consistent from the periphery to the IC, but sharper in A1. This may be due to a number of reasons; the assumptions we make that allow us to interpret the activity of cells and convert firing rates into signal detection performance may not hold for the more complex A1 cells, or perhaps cortex uses a sub-optimal method for pooling cell activity when performing a detection task. The cause is still being investigated.

Electrophysiological Recordings from the Parabelt Areas in Macaques
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Background
According to anatomical connection patterns, primate auditory cortex has 3 hierarchical regions - core, belt and parabelt. Every region contains several areas. In macaques, the core and most of the belt regions stretch posterior-to-anterior on the superior temporal plane. Tonotopic gradients across areas run parallel in core and belt areas. Parabelt is elongated on the superior temporal gyrus (STG) and divided into caudal (CPB) and rostral (RPB) areas. Based on connection patterns and functional features, CPB and RPB are speculated to be parts of caudal "where" and rostral "what" pathways, respectively. Functional segregation between areas was suggested for the macaque belt by electrophysiological studies. However, the functions and physiological properties of macaque STG remain unknown, as does the relationship between
CPB and RPB. We mapped auditory responses in PB of an awake macaque monkey, using a chronic chamber technique to make electrode penetrations perpendicular to STG.

Methods
Linear array multielectrodes were used to record field potential and multiunit activity (MUA) simultaneously across the laminar depth of STG cortices. We used a battery of sounds including pure tones (0.3 to 32 kHz), 1/3 oct. band-pass noises (BPN), broad band noises (BBN), click trains, sinusoidal amplitude modulation (SAM) tones, FM sweeps, and conspecific vocal sounds, and were delivered binaurally. We defined a best frequency (BF) at each penetration site using responses to tones and BPN.

Results
Many sites in STG appeared “tuned” to tones and BPN. Topographic patterns of BF were similar between tones and BPN, with a common low BF zone that could be a border between CPB and RPB. In the putative CPB, most sites responded to tones and formed a tonotopic gradient roughly in parallel to those in A1/CL. About 50% of RPB sites did not respond to tones, and RPB had no clear tonotopic gradient. Latencies of responses to BBN were longer in RPB than CPB.

Conclusion
Our implant technique allowed direct access to Macaque STG. We mapped acoustic response properties in parabelt areas. While PB areas may prefer complex or vocal sounds to simple sounds, they also responded surprisingly well to tones and BPN with spectral tuning. RPB had smaller fraction of tone-responsive sites, suggested more selectivity to sounds in RPB than CPB. Shorter response latencies in CPB suggested a possibility of CPB-to-RPB projections.

Impact of Low Dose GABA on Temporal Responses: Comparison between MGB and IC
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Background
Previous studies demonstrated that neurons at higher levels of the auditory system respond to sinusoidal amplitude modulated (SAM) stimuli with large and complex variations in spike rate. Our preliminary medial geniculate body (MGB) findings and a number of earlier studies in the inferior colliculus (IC) found that GABA played a role in controlling response rate and shaping SAM response properties. The presence of high affinity GABAA receptors in MGB and the present iontophoretic studies suggest MGB neurons display a heightened sensitivity to GABA application. The present study quantifies the potency of GABA under parallel experimental conditions comparing the GABA 50% inhibitory dose (ID50) for MGB and IC neurons.

Methods
A commercially available, carbostar-6, was used for recording and drug delivery (GABA, 10mM). Rate modulation transform functions (rMTFs) were generated from responses of isolated single units and occasional small clusters within MGB and IC in response to randomly presented SAM (2-512 Hz, 450ms duration, 100% modulation depth, BBN or BF carrier). Retaining currents were set at -15nA and drug delivery dosage (eject current amplitude) ranged from 0nA up to 100nA with 5 or 10nA steps.

Results
In MGB and IC, rMTFs can be categorized into low pass, high pass, bandpass, band-reject response types. GABA dose-dependent effects were obtained from 34 MGB and 10 IC units, focused on bandpass and band-reject response types. Spike counts at -15nA retaining current was considered as the control condition for each unit. The percentage change in spike count was obtained at different ejection doses. A linear fit curve was applied to the percentage value against ejection current for each unit and ID50 values were calculated. The median ID50 in MGB was 11.72, and the mean was 14.01 ± 3.00 (Mean ± SE). IC neurons were significantly less sensitive GABA application than MGB neurons (p < 0.001, independent t-test), with a median ID50 value of 51.28 and mean of 49.56 ± 6.75.

Conclusion
These results find that MGB neurons are three times as sensitive to GABA application as IC neurons. This difference is likely due to the presence of high affinity GABAA receptors in MGB. It has been suggested that, due to the presence of these high affinity GABAAARs, ambient GABA levels may uniquely gate and shape the ascending information from MGB to auditory cortex. These data also suggest a specific role for these receptors in temporal processing in the MGB.
Cortical spectrotemporal processing with age-related hearing loss
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Background
Presbycusis (age-related hearing loss) is the most prevalent hearing impairment in humans. It affects ~40% of people older than 65 years. One major symptom of presbycusis is impaired speech recognition. Hearing aid amplification is only a partial solution, indicating central auditory system plasticity. The central changes have been insufficiently characterized. Here we studied cortical processing of frequency modulated (FM) sweeps to evaluate spectrotemporal representation plasticity in a mouse model of presbycusis (C57bl/6 strain).

Methods
Single unit electrophysiology in mice anesthetized with ketamine/xylazine was used to record FM selectivity. We compared cortical (both A1 and AAF) FM sweep rate selectivity across three age groups: Young (‘Y’, 1-2 mo), Middle-age (‘M’, 6-8 mo) and Old (‘O’, 14-20 mo). Sweep rates between 0.06-22 kHz/msec (linear sweeps) were tested. Neurons were classified as fast-pass, slow-pass, band-pass or all-pass based on response selectivity.

Results
Three main results were observed with age: 1) There were less fast-pass and band-pass neurons and more slow-pass neurons. 2) Selectivity of fast-pass and band-pass neurons was shifted towards slower sweep rates. Overall selectivity of band-pass neurons was reduced 3) Response variability to stimulus repetitions in terms of magnitude, first-spike latency and interspike intervals was increased.

Conclusion
Thus, processing of spectrotemporal cues becomes slow and noisy with presbycusis.

Consonance-dependent Modulation of Phase Synchrony in the Auditory Cortex of Rats
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Background
Many sounds in the environment have a rich sound spectrum, which is related to the qualia of the sound such as consonance or dissonance of chords. However, the information processing of the consonance of the chord in the auditory cortex is not fully understood. In this study, we targeted a phase locking value (PLV) of local field potential (LFP) and investigated whether PLV represents consonance of the chord consisting of two pure tones.

Methods
A microelectrode array with a grid of 96 recording sites recorded LFPs in the fourth layer of the auditory cortex of anesthetized naive rats in response to continuous pure tones and chords. In the pure tone condition, we randomly presented 9 pure tones (12, 13.5, 14.4, 15, 16, 19.2, 20, and 24 kHz, 60 dB SPL) for 30 seconds continuously, each of which was interleaved with a silent block of 30 seconds. In the chord condition, we made 8 chords by combining 12-kHz pure tone and one of other 8 pure tones, and presented them in the same way as the pure tone condition. The recorded LFPs were bandpass filtered in 5 bands (theta, 4 - 8 Hz; alpha, 8 - 14 Hz; beta, 14 - 30 Hz; low gamma, 30 - 40 Hz; high gamma, 60 - 80 Hz) and PLVs were calculated in each band. First, a median of PLV among all pairs of recording sites within the auditory cortex in each silent and stimulus block was calculated. Second, a difference of median of PLV in the stimulus block with respect to that in the adjacent silent block was calculated as \(\Delta PLV\). Finally, a difference of \(\Delta PLV\) of a chord with respect to \(\Delta PLV\) of the higher-frequency composition tone of the chord was evaluated in consonant (low composition, 12 kHz; high composition, 15, 16, 18 or 20 kHz) and dissonant chords (low composition, 12 kHz; high composition, 13.5, 14.4 or 19.2 kHz) (Fig.1).

Results
\(\Delta PLVs\) of consonant chord were significantly larger than \(\Delta PLVs\) of pure tone in all bands, while \(\Delta PLVs\) of dissonant chord were not significantly larger except alpha band (Fig. 2). Especially in beta, low gamma and high gamma band, \(\Delta PLV\) increase in a chord was larger in consonance than dissonance.

Conclusion
These results suggest that the phase synchrony within the auditory cortex is modulated according to the complex sound spectrum, such as consonance of the chords.
Measurement of Auditory Cortical Network Responses to Potassium Channel Openers

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Background
Potassium (K⁺) channels are primary regulators of neuronal excitability, with significant roles in neuropathic pain, epilepsy, cardiac arrhythmia, hearing loss, and tinnitus. In recent years, a novel class of drugs – K⁺ channel openers – has been shown to exert anticonvulsive and neuroprotective effects by activating K⁺ currents that induce hyperpolarization. Retigabine, a first-in-class channel opener of a voltage-gated K⁺ channel subclass (Kv7), is the prime candidate currently undergoing clinical trials for the treatment of epilepsy.

Methods
We present quantitative measurements of the modulatory effect of K⁺ channel openers on the activity pattern of spontaneously-active neuronal networks, and discuss the potential therapeutic value of K⁺ channel openers on pathological auditory cortical activity such as tinnitus. Networks derived from dissociated auditory cortices of mouse embryos were grown on microelectrode arrays (MEAs) to assay the drug-induced changes in network spike production, burst patterns, and action potential waveforms.

Results
We assessed the effects of retigabine, flupirtine (structural analog of retigabine), isopimaric acid, and NS1619 (activator of large-conductance calcium-sensitive potassium channels). The EC₅₀ values were 8.0, 4.0, 7.8 and 5.8 µM, respectively. Despite comparable potency – flupirtine (EC₅₀: 4.0 µM) > NS1619 (5.8) > isopimaric acid (7.8) > retigabine (8.0) – and full efficacy, each K⁺ channel opener exhibited differential burst properties under low concentration (2.0 µM), and distinct K⁺ waveform component alteration of action potentials under high concentration (20 µM).

Conclusion
This study, using the in vitro auditory cortical network model coupled with the MEA platform for rapid drug screening and identification of mechanisms, may serve as the basis for developing and re-purposing K⁺ channel openers as potential drugs for the treatment of tinnitus.

Multiscale Functional Imaging of Auditory Cortical Areas in Awake Mice

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Background
Classic electrode studies support robust spatial organization of sensory cortices, comprised of distinct areas with remarkable functional specialization and segregation. In the mouse auditory cortex, auditory subregions with distinctive receptive field properties have been observed: AI, AAF, UF, DP, and AII. For example, in AI and AAF, preferred frequencies appear organized spatially along a rostrocaudal gradient; neurons in UF prefer high frequencies (>40 kHz); and AII contains neurons with multi-peaked tuning (J Comp Physiol A., 181(6):559-71).

However, these studies may underestimate the actual heterogeneity of neuronal responses because the methodologies used may be biased towards strongly responding neurons. More recent two-photon Ca²⁺ imaging studies, where a large number of individual neurons can be imaged simultaneously, have revealed diverse receptive fields among nearby
neurons (Nat Neurosci., 13(3):353-68) and even between nearby spines on the same neuron (J Neurosci., 32(5):1589-601). These results suggest a high diversity of response properties in auditory cortex, contrasting with the well-organized and distinctive auditory field maps expected from prior studies. To reconcile these contrasting perspectives, we here combined functional transcranial fluorescence imaging registered with two photon Ca\(^{2+}\) imaging of individual neurons in the auditory cortices of awake mice.

**Methods**

The left auditory cortical areas of adult mice were initially characterized by functional transcranial fluorescence imaging. The large imaging field (~5 mm) permitted simultaneous imaging of all auditory fields. Next, we undertook two-photon Ca\(^{2+}\) imaging of layer 2/3 neurons, as registered to transcranial images via blood vessel landmarks.

**Results**

Transcranial imaging rapidly identified multiple, well-demarcated auditory cortical areas, including AI, AAF, UF, and All. Response properties and spatial organization exhibited remarkable agreement with the consensus view of earlier electrode studies, such as strong rostrocaudal gradients of preferred frequency in AI and AAF. Under two-photon Ca\(^{2+}\) imaging, the average response, taken by summing firing across individual neurons, accorded well with the global profile of corresponding cortical regions. Beyond this clear overall congruence, single-neuron activity exhibited significant divergence in precise responses among neighboring neurons, leaving room for higher-order representations of acoustic stimuli. Compared to anesthetized mice, single-neuron activities in awake animals were often enhanced, favoring well-resolved characterization of response profiles.

**Conclusion**

The combination of transcranial functional imaging and two-photon Ca\(^{2+}\) imaging promises to reveal how global receptive-field organization intersects the neural activity of individual neurons across multiple spatial scales.

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**Morphological analysis of auditory thalamus in BXSB/MpJ mice**

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**Background**

In the BXSB/MpJ inbred mouse strain, approximately half the animals spontaneously develop nests of displaced neurons in cortical layer I, called ectopias. Neocortical ectopias have also been reported to occur in humans with developmental disorders such as dyslexia. Interestingly, although the ectopias occur outside the auditory cortex, both human studies and studies of rats with induced cortical malformations have observed an association with anatomical abnormalities within the auditory thalamus. Previously, we have shown that ectopic male BXSB/MpJ mice have an auditory thalamic processing deficit, despite apparently normal hearing sensitivity. The aim of this current study was to determine whether these animals also have anatomical abnormalities in the auditory thalamus.

**Methods**

Auditory brainstem responses and extracellular thalamic recordings were collected from urethane-anaesthetised male BXSB/MpJ mice during monaural presentations of free-field auditory stimuli. All experiments were necessarily performed blind to the ectopic status of the animal. Following in vivo physiological recordings, brain tissue was processed for histology. Alternate coronal brain sections were stained for cytochrome oxidase to differentiate thalamic subdivisions, and Nissl substance to reveal cortical ectopia and thalamic cell density. Volumes of cortical ectopias and auditory thalamic subdivisions were calculated using area estimates obtained by drawing borders on stained sections. Cell packing densities were estimated with a pixel-based density index, calculated from the mean of median pixel intensities across different image squares for the same subdivision and animal.

**Results**

Most ectopic animals had a single ectopia (volume 0.001-0.036 mm\(^3\)), located in the motor cortex. Ectopia volume was found to correlate with auditory thalamic processing deficits; animals with larger ectopia in motor cortex had larger deficits (p < 0.05). However, anatomical analysis showed no significant differences in thalamic volume or cell density between ectopic and non-ectopic mice, for either the whole auditory thalamus or any individual subdivision.

**Conclusion**

Cortical ectopias in BXSB/MpJ mice were not associated with any significant changes in auditory thalamic morphology; however, ectopia volume correlated with physiological deficits in auditory thalamic processing. These results suggest that
the deficits might arise outside the auditory thalamus, and perhaps that motor cortex ectopias and auditory thalamic processing deficits have a common underlying developmental cause.

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An emergent spectro-temporal representation for natural sounds based on sustained firing of central auditory neurons
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Background
Natural sounds evolve over many concurrent timescales, from tens to hundreds of milliseconds, yet the mechanisms by which the auditory system calibrates internal neural dynamics to the information-bearing components of these signals remains unclear. Recent data suggests the presence of a neural code based on sustained firing rates, whereby central auditory neurons exhibit strong, persistent responses to their preferred stimuli. In this work, we examine how this code may underlie aspects of how the auditory system learns to represent sound.

Methods
We explore how a coding strategy that maximizes sustained firing rates of model neurons influences the shapes of spectro-temporal receptive fields (STRFs) in the central auditory system.

Results
We demonstrate the emergence of richly structured STRFs that capture the structure of natural sounds over a wide range of timescales, and show how the emergent ensembles resemble those commonly reported in physiological studies. We also show how the emergent ensembles capture the full range of spectro-temporal modulations observed in natural sounds, forming a discriminative representation that captures the full range of modulation statistics that characterize natural sound ensembles.

Conclusion
The results suggest that a coding strategy based on sustained firing rates may underlie part of the neural code for representing sounds in the central auditory system.
Cell type-specific cholinergic regulation of action potentials in layer 6 neurons of auditory cortex

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Background
Layer 6 of primary auditory cortex (A1) contains neurons that control cortical gain of auditory information by providing feedback to primary thalamic neurons in the ventral division of medial geniculate nucleus (MGv) and by modulating thalamorecipient neurons in the middle layer of A1. Since cholinergic inputs to A1 regulate receptive field of MGv neurons, neuromodulatory mechanisms of cortical neurons, particularly of layer 6 neurons, may be critical for gating of auditory information. Many previous studies have revealed cholinergic regulation of neuronal excitability in the neocortex. However, little has been studied about the regulation of layer 6 neurons. As a step toward understanding cellular mechanisms of sensory gating, we investigate intrinsic membrane properties and cholinergic regulation of neuronal excitability in layer 6 neurons.

Methods
In auditory thalamocortical slices, we have recorded membrane potentials of layer 6 neurons using whole-cell patch clamp recording techniques. Cholinergic ligands were tonically applied to examine their effects on intrinsic membrane properties and action potentials elicited by current injections. Cell type and morphology of recorded neurons were analyzed by examining the expression of glutamic acid decarboxylase 67 kDa (GAD67) - an inhibitory neuronal marker - using immunofluorescent staining method and by histochemical analysis of biocytin-filled neurons.

Results
Based on spike firing properties and immunofluorescent staining, we divided neurons into three groups: regular spiking (RS), inhibitory RS (iRS), and fast spiking (FS) neurons. Tonic application of acetylcholine (ACh; <100 μM, ~5 min) increased resting membrane potentials, input resistance, and spike frequency while decreasing spike onset time in all layer 6 neurons recorded. In RS neurons, ACh decreased spike threshold for the first spike as well as repolarizing afterhyperpolarization (fAHP) with little changes in spike width and peak amplitude. In iRS neurons, ACh greatly increased spike threshold, spike width, and fAHP while decreasing peak amplitude. Meanwhile, in FS neurons, ACh had little effects on spike threshold, peak amplitude, and spike half-width, but decreased fAHP.

Conclusion
ACh increased the excitability of all layer 6 neurons we recorded by increasing resting membrane potentials and input resistance. Analysis of action potentials revealed the difference in cellular mechanisms of cholinergic regulation in RS, iRS, and FS neurons.
Neurochemical, physiological and anatomical studies of the auditory thalamotectal projections of the mouse
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Background
Descending projections are found at virtually every level of the central auditory system. Though many of these projections have garnered attention in recent years, the auditory thalamotectal pathway remains poorly characterized. The purpose of the current study was to provide an initial characterization of this pathway, using neurochemical and physiological approaches. We examined immunoreactivity for two calcium-binding proteins, calbindin and calretinin, in auditory thalamotectal cells since these proteins are found in the medial and dorsal divisions of the medial geniculate body. As thalamotectal cells arise from the dorsal and medial divisions of the medial geniculate body, we hypothesize that they will be immunoreactive for either calbindin or calretinin, like their neighboring cells in these regions. We also studied the electrophysiological properties of identified thalamotectal cells.

Methods
A fluorescent retrograde tracer, either red polystyrene Fluospheres (for physiological studies) or Fluorogold (for immunostaining studies), was injected into the inferior colliculus of mice to label thalamotectal cells. Fluorogold-injected animals were perfused, had brains removed and tissue sections that contained labeled thalamotectal cells were immunostained for either calbindin or calretinin. Living brain slices were obtained from Fluosphere-injected animals, and whole-cell patch clamp was used to study the physiological properties of these cells. Recorded cells were filled with biocytin for morphological characterization.

Results
We found that the vast majority of retrogradely-labeled cells (>90%) were negative for either calbindin or calretinin. Positive controls with retrogradely labeled thalamocortical cells showed strong immunostaining for both calbindin and calretinin. Physiologically, thalamotectal cells showed little bursting behavior and did not show typical bitufted thalamic morphology.

Conclusion
Mouse auditory thalamotectal neurons do not exhibit strong positivity for calbindin or calretinin, do not show bursting and do not have typical thalamocortical cell morphology. Based on these results, it is possible that these cells form a distinct class of thalamic neurons that differ from neighboring auditory thalamic cells.

Temporal Modulation Context Affects the Cortical Representation of Amplitude Modulated Signals in Awake Monkeys
Ralph Beitel1, Brian Malone1, Maike Vollmer2, Marc Heiser3, Christoph Schreiner1
1UCSF, 2Comprehensive Hearing Center Wuerzburg, Germany, 3UCLA

Background
Sinusoidal amplitude modulated (SAM) signals are often used in psychophysical studies of temporal processing in the auditory system. Proposals about encoding of SAM sounds in the brain include filterbank models tuned to modulation frequencies as well as models that favor information processing across modulation frequencies. In this physiological study we presented pairs of modulated stimuli (SAM1/SAM2) to awake monkeys and investigated the effects that SAM1 had on cortical responses to SAM2.

Methods
Two squirrel monkeys were trained to sit facing a speaker with head restrained during recording sessions. Multiunit responses were recorded in core auditory cortex with a linear 16-channel silicon probe. The carriers for modulated signals were pure tones near channel best frequency (BF) or 2-octave bandwidth noise centered on channel BF. SAM2 modulation frequencies were typically presented from 4 to 512 Hz in pseudorandom order. The effects of stimulus context were investigated by comparing the shapes of modulation transfer functions (MTFs) when each SAM2 stimulus (1-s) was preceded immediately by a SAM1 stimulus (1-s) at a constant frequency (4, 10, 32, or 96 Hz) or by an unmodulated stimulus. Responses to SAM2 signals presented alone were also studied. Modulation depth was 100% for all SAM stimuli.

Results
The spike rates to SAM2 presented alone or preceded by an unmodulated stimulus were similar and were usually higher than spike rates to SAM2 stimuli preceded by SAM1 stimuli. In comparing the effect of different SAM1 modulation
frequencies on SAM2 spike rates, higher SAM1 modulation frequencies generally resulted in a significant reduction in SAM2 spike rates when compared to lower SAM1 frequencies. This reduction in spike rates occurred over a broad range of SAM2 modulation frequencies (± 2 oct) and tended to be larger for SAM2 modulation frequencies nearer to the modulation frequency of the preceding SAM1 segment. Compared to rate MTFs, the effects of stimulus context on MTFs defined by temporal metrics (vector strength and trial-to-trial response similarity) were small.

**Conclusion**
Temporal context significantly affects cortical responses to SAM signals in a manner consistent with the idea of adaptation within a broadly tuned modulation frequency channel. In contrast, a preceding unmodulated stimulus had no consistent effect on the responses to modulated stimuli, suggesting that preceding modulation was necessary as well as effective for the observed spike rate effects.

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**Noradrenergic gating of long-term cortical synaptic plasticity**

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**Background**
Neuronal networks of the cerebral cortex are plastic, maintaining the capacity to reorganize throughout life. While neuromodulator release is required for cortical plasticity, it is uncertain how subcortical neuromodulatory systems, such as the noradrenergic locus coeruleus, interact with and refine cortical circuits.

**Methods**
Here we determine the dynamics of auditory cortical receptive field plasticity at the synaptic and spiking levels using in vivo whole-cell recording. Adult rats were anesthetized, stimulation electrodes were implanted in the locus coeruleus, and a craniotomy performed over the primary auditory cortex. After mapping the tonotopic field of the primary auditory cortex, intracellular recordings were made (usually in layer V) from cortical neurons, and pure tones of varying frequencies and intensities were presented to the animal to characterize tonal receptive fields.

**Results**
Pairing sensory stimulation (pure tones) with locus coeruleus activation (to release noradrenalin) dramatically changed the tuning properties of cortical neurons. For most recordings, pairing induced large increases of tone-evoked synaptic and spiking responses, degrading neuronal tuning across all frequencies. Changes in tuning curve shape continued to evolve over tens of minutes to emphasize and disambiguate the paired stimulus from other unpaired stimuli. Multiple cell recordings from the same animal for hours after a single episode of locus coeruleus pairing suggested that these changes in AI tuning stabilized after three to six hours and could persist for 11+ hours.

Furthermore, some AI neurons were initially unresponsive to auditory stimuli. During and after locus coeruleus pairing, however, tone-evoked responses could suddenly emerge, persisting for the duration of post-pairing recordings (30+ min). This suggests the involvement of the locus coeruleus arousal system in the integration of ‘silent neurons’ into a specific auditory memory trace or cortical representation of certain, potentially behaviorally important sensory stimuli.

Moreover, our data suggests that the locus coeruleus itself seems to undergo local activity changes that at least partially underlie the cortical changes observed.

**Conclusion**
We hypothesize that such changes to cortical tuning curves have important implications for the detection and/or discrimination of different sensory inputs, and that the subcortical sensitization of locus coeruleus plays a major role in the cortical changes observed.
The effect of sound source location on multi-peak frequency tuning in the primary auditory cortex of marmoset monkeys
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Background
When the listening environment contains multiple sound sources emitting sounds from various spatial locations, sound source identification becomes difficult to achieve. On the other hand, the auditory system routinely streams together isolated sound elements into a coherent perceptual image using multiple grouping cues, such as harmonicity and common onset. For example, pure tones that are harmonically related can be fused into a single auditory event regardless of their spatial locations. To understand the underlying neural mechanisms, this study investigated whether auditory cortex neurons treat sounds originating from the same source or different sources in different ways.

Methods
Single-unit data was collected from the primary auditory cortex (A1) of awake marmoset monkeys. The experiments were conducted in free field with multiple loudspeakers positioned in the frontal hemifield. To analyze the efficacy of spectral integration in A1, our analyses focused on neurons showing multi-peak frequency tuning profiles in response to a pair of pure-tone stimuli delivered from the same loudspeaker.

Results
The two-tone paradigm revealed substantial inhibitory activity in frequency tuning of neurons. We observed that many multi-peak neurons showed either excitatory or inhibitory sensitivity at distant frequencies with particular harmonic ratios to their best frequency (BF), e.g., 1/2BF or 2BF. Interestingly, presenting the two tones from two different spatial locations did not change the multiple-peak tuning profiles of a majority of neurons. In some neurons, inhibitory interactions were observed when a pure tone with a distant frequency (e.g., 2BF) was presented from the location showing no excitatory responses to BF.

Conclusion
These observations provide an interesting example of location invariance in cortical spectral processing. The data show that A1 neurons can integrate harmonically related excitatory or inhibitory inputs across a broad spatial domain.
Structured Sparse Encoding for Automated Segmentation of Calcium Imaging Data
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¹Johns Hopkins University

Background
Advances in fluorescent microscopy and improvements in synthetic and genetically encoded calcium indicators have enabled recordings of calcium dynamics from large numbers of neurons. However, as the number of recorded neurons increases, segmenting the neuron boundaries from the raw fluorescent data becomes increasingly challenging and presents a significant bottleneck in the data analysis. Several methods have been suggested to automate the image segmentation process but their application has been limited by decreased performance in the presence of image noise, overly restrictive model assumptions, or excessive computation time.

Methods
To address these issues, we have developed an image segmentation algorithm which uses structured sparse encoding to learn a basis of dictionary elements and then decomposes the fluorescence data into segmented regions and estimated spike trains simultaneously. To demonstrate the utility of our method we use it to segment two photon calcium image data taken from the auditory cortex of awake mice. The cortex was virally transfected with the genetically encoded calcium indicator GCaMP5, and images were acquired at multiple levels of optical resolution. Robust activity was observed in response to both natural mouse calls and pure tone stimuli.

Results
Our method requires minimal user input or parameter tuning, makes no assumptions about region morphology, is robust to noise, and is easily modified to accommodate a wide variety of calcium dynamics. When applied to data taken at multiple levels of optical resolution, our method is able to identify neural boundaries and estimate spike trains from both cell bodies and dendritic processes.

Conclusion
Structured sparse encoding provides a robust, efficient, and highly flexible method to decompose neural calcium image data into both segmented cell boundaries and estimated spike trains.
**Intracortical Inputs to Neurons in Layer 4 of Primary Auditory Cortex**

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**Background**

One task of the auditory system is to identify auditory objects in the environment. Auditory objects often contain component frequencies from across the hearing range. This presents unique challenges for the neural circuitry that processes sound, as it requires extensive lateral connectivity between different parts of the tonotopic map. One of the earliest sites in the auditory pathway where convergence from across the entire cochlear partition can take place (aside from octopus cells of the cochlear nucleus) is the primary auditory cortex (A1). Here, we investigate the sources of intracortical input from across the tonotopic axis onto to Layer (L) 4 neurons in mouse primary auditory cortex.

**Methods**

We use thalamocortical slices of P35-43 CBA/CaJ mouse A1 that cut across the tonotopic map. We then record excitatory post-synaptic currents from L4 neurons while stimulating potential presynaptic partners throughout A1 using laser scanning photostimulation and glutamate uncaging. Uncaging sites were designed on a dense, overlapping hexagonal grid so that any presynaptic neuron will likely be activated by stimulation at multiple adjacent sites. This approach allows us to use the spatial correlation of stimulation-evoked events to more confidently determine the presence and location of presynaptic cells.

**Results**

While input maps to individual L4 neurons are heterogeneous, they share some common features. Discrete, discontinuous inputs from other L4 neurons ~300 μm caudal (towards the low-frequency end of the tonotopic map) to the cell are evident in about 40% of cells, as are similar inputs from L4 neurons ~300 μm rostral. Inputs were also seen from L6 neurons both ~300 μm rostral and caudal, although few cells receive inputs from all of these locations. L4 neurons also receive “columnar” inputs. Nearly all cells receive L4 input from the immediate vicinity of the cell, while approximately 30%, 22% and 50% of cells receive same-column input from L2/3, L5, and L6, respectively.

**Conclusion**

These results suggest that information from different tonotopic locations is combined with thalamocortical inputs in L4 neurons. The non-contiguous but regular organization of the input zones is consistent with a modular processing circuit. The (putative) cross-tonotopic connections shown here may be tuned to help amplify or fill-in harmonic structure, aiding our ability to detect, identify, and attend to auditory objects.

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Cortical Auditory Evoked Potentials to Frequency Changes with Varied Size, Velocity and Direction
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Background
Cortical auditory evoked potentials (CAEPs) in response to brief changes in sounds (e.g., vowel transitions, or changes in level or frequency) may be suitable for assessing suprathreshold neural processes related to speech perception. These CAEPs evoked by sound changes are also referred to as acoustic change complexes (ACCs) and are composed of a typical P1-N1-P2 waveform morphology (N1 latency ≈ 100 ms after stimulus change). In this study we explored ACCs to small frequency modulation (FM) sweeps following 1000-Hz pure tones. Size, velocity and direction of the FM sweep were systematically varied.

Methods
Ten young adult human subjects with normal hearing participated. Stimuli were tones of 3300 ms with three components: a) a reference tone of 1000 Hz with a duration of 3000 ms, b) an FM sweep with a frequency change Δf, c) a tone with a frequency of 1000-Δf or 1000+Δf Hz and a duration of 300 ms. The silent interval between stimuli was 200 ms. Sound level was 75 dB SPL. Three frequency changes were used: 0.029, 0.086 and 0.257 octave; four FM velocities were used: 3.2, 9.5, 28.6 and 85.7 octave/s; both up and down sweep directions were presented. The ACCs were recorded with Ag/AgCl electrodes at Cz referenced to the contralateral mastoid. For each stimulus condition, ACC waveforms were averaged over 100 presentations.

Results
As previously shown in literature, the ACC amplitude increased with increasing frequency change. The amplitude doubled when the change was increased with a factor 9. Further, the amplitude increased with about 25% when velocity was increased with a factor 27. Finally, the ACC amplitudes tended to be larger for downward frequency changes.

Conclusion
The primary factor determining the ACC amplitude is the size of the frequency change. A secondary factor is velocity of the change. For clinical purposes, for instance when assessing thresholds of responses to frequency changes, an appropriate stimulus to evoke ACCs would be a fast FM sweep following a long pure tone.

Subcollicular sources of projections to the auditory thalamus in the rat
Enrique Saldaña¹,², F.-Javier Bernardo¹, Yongming Jin³, Marcelo Gómez-Álvarez², M.-Auxiliadora Aparicio¹, Emmanuel Márquez¹, David M. Sloan³, Albert S. Berrebi³
¹University of Salamanca, ²Institute of Biomedical Research of Salamanca (IBSAL), Spain, ³West Virginia University School of Medicine

Background
It is generally assumed that the main source of ascending projections to the auditory thalamus is the inferior colliculus (IC). However, several isolated reports suggest that the medial geniculate body (MGB) receives direct projections from different auditory subcollicular nuclei (e.g. Henkel, Brain Res 259:21–30, 1983 [cat]; Angelucci et al., J Comp Neurol 400:417–439, 1998 [ferret]; Malmierca et al., J Neurosci 22:10891–10897, 2002 [rat]; Berrebi et al., ARO Abstr 35:250, 2012 [rat]). This is particularly relevant because evidence presented at this meeting indicates that, contrary to the long-held tenet, the medial division of the MGB (MGBm) is not a major target of the IC (Saldaña, Symposium on “The non-lemniscal auditory thalamus”).

Methods
To systematically investigate the sources of ascending projections to the auditory thalamus, we made iontophoretic injections of the sensitive retrograde tracer FluoroGold into the auditory thalamus and analyzed the distribution of the neuronal cells bodies labeled in all subcollicular auditory nuclei.

Results
We obtained cases with injection sites centered in different regions of the auditory thalamus, including the ventral, dorsal and medial divisions of the MGB, the posterior intralaminar nucleus, the marginal zone and the posterior limitans nucleus. In the classical auditory nuclei, abundant retrogradely labeled neurons were found in the contralateral dorsal cochlear nucleus and lateral superior olive; in the ipsilateral medial superior olive (MSO), superior paralaminar nucleus (SPON), and ventral nucleus of the lateral lemniscus; and in the dorsal nucleus of the lateral lemniscus and the nucleus sagulum of
both sides. We also found numerous labeled neurons ipsilaterally in the ventrolateral tegmental area, located medial to
the ventromedial portion of the VNLL, as well as in an ill-defined territory wedged between the medial nucleus of the
trapezoid body and the SPON. In the cases with the most retrograde labeling, the percentage of labeled neurons in some
nuclei, such as the ipsilateral MSO and SPON, approached 50%.
Because FluoroGold injection sites tend to be large and usually include more than one subdivision of the auditory
thalamus, we are currently performing a quantitative, multivariate analysis to infer the relative contribution of each one of
the subcollicular nuclei to the innervation of each region of the auditory thalamus.

Conclusion
The auditory thalamus is innervated by numerous neurons in most subcollicular auditory centers. These findings improve
our understanding of the auditory pathway and provide novel morphological frameworks for future functional studies.

Oxytocin-based neuromodulation of mammalian social behavior
Bianca Jones1, Robert Froemke2
1New York University School of Medicine, 2New York University
Background
Social interactions are essential for normative development and lifelong physical and mental health. The neuromodulator
oxytocin is believed to be important for some forms of social behavior, specifically parent-child bonding and maternal
behavior. Here we investigate a fundamental form of rodent social behavior that depends on both social cues and
modification of neural substrates: pup retrieval.

Methods
To initiate pup retrieval behavior, mouse pups produce ultrasonic vocalizations (USVs) when separated from the nest.
Dams and other experienced caregivers use these acoustic signals to locate and retrieve isolated pups. Importantly, virgin
female mice can also learn a variety of maternal behaviors, including pup retrieval, and we hypothesize that oxytocin
plays a major role in this form of learned social behavior (Pedersen et al., Science 1982). However, it is unclear how
oxytocin impacts the activity of the cortex, leading to the robust increase in maternal behavior.

Results
We confirmed that virgins can also learn to retrieve pups. Furthermore, learning is accelerated after either systemic or
cortical application of oxytocin. Virgins were first tested for initial retrieval abilities. Virgins were then co-housed with a
dam and her pups for up to three days, with some animals receiving oxytocin and other animals receiving saline. After 12
hours of co-housing, 16/31 (~50%) animals injected with oxytocin learned to retrieve pups, but only 6/24 (25%) control
saline-injected virgins retrieved pups. Similarly, 7/9 animals receiving oxytocin infusion into left auditory cortex via cannula
retrieved pups within 12 hours, but 0/4 cannulated virgins receiving saline infusion retrieved pups. Finally, infusing the
GABA-A receptor agonist muscimol into left auditory cortex, to inactivate local circuitry, reduced or fully prevented pup
retrieval in 6/9 experienced animals. Infusion into right auditory cortex did not seem to significantly affect retrieval.

Conclusion
This demonstrates that left auditory cortex is required for USVs to engage pup retrieval. Furthermore, these data suggest
that oxytocin-based neuromodulation, paired with acoustic stimuli, can modify auditory cortex to enable or improve
learned social behavior.
Functional Organization of Spectrotemporal Processing in Human Temporal Lobe.
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Background
Functional organization within the nervous system often reveals key insights into the information being represented and the processing taking place. In the human auditory system, tonotopic organization is a prominent organizing principle up through the primary auditory cortex, but less is known about the functional organization of higher auditory areas. It is of considerable interest to determine what organizing principles, if any, are present in higher-order auditory areas in humans. Here we use ECoG recordings in conjunction with advanced techniques for computing spectrotemporal receptive fields to investigate what forms of functional organization are present in areas beyond primary auditory cortex.

Methods
To characterize functional organization in higher-order auditory areas, we recorded auditory responses to natural speech in the temporal lobe of awake humans using electrocorticography (ECoG). Although restricted to rare clinical settings, ECoG provides high spatiotemporal resolution of neural activity, which is ideal for investigating questions of functional organization in the auditory system. Using the high gamma component of ECoG recordings in three patients who passively listened to speech, we computed spectrotemporal receptive fields (STRF’s) using maximally informative dimension (MID) analysis and looked for functional organization based on the spatial distribution of STRF’s across the temporal lobe.

Results
Parameters derived from each STRF, including best spectral modulation, best temporal modulation, best frequency, bandwidth, and temporal integration were mapped onto cortex to characterize the functional organization of spectrotemporal processing. One prominent outcome of this analysis shows functional organization based on spectral and temporal modulation tuning. First, there is a spectral-temporal trade-off in which regions tuned to high spectral modulations prefer low temporal modulations while regions tuned to high temporal modulations prefer low spectral modulations. Second, there are smooth transitions from high spectral modulation tuned regions to high temporal modulation tuned regions thus forming an organized distribution of modulation tuning across temporal lobe auditory cortex.

Conclusion
These results indicate that modulation tuning is a dominant organizing feature within temporal lobe auditory cortex. Although other forms of functional organization are present, organization based on modulation tuning is most descriptive of temporal lobe activity in response to complex signals like speech. At the level of the temporal lobe, these maps imply the extraction of spectrotemporal modulation content is a critical step in the process of speech perception.
Functional topography of thalamocortical and intracortical inputs to layers 4 and 6b
Charles Lee¹, Kazuo Imaizumi¹
¹Louisiana State Univ. School of Veterinary Medicine

Background
In the auditory forebrain, thalamocortical axons transfer information from the thalamus to layers 4 and 6b of sensory cortical areas. Yet, receptive field properties in layer 6 are generally more complex than those in layer 4. Thus, it remains an open question whether such differences reflect distinct inheritance patterns from the thalamus or if instead they are derived from local cortical circuits.

Methods
To distinguish between these possibilities, we utilized in vitro slice preparations containing the intact thalamocortical pathways in the auditory and somatosensory systems. Responses from neurons in layers 4 and 6b that resided in the same column were recorded using whole-cell patch clamp. Laser-scanning photostimulation via uncaging of glutamate in the thalamus and cortex was then used to map the functional topography of both thalamocortical and intracortical inputs.

Results
We found that the thalamocortical inputs to layers 4 and 6b originated from the same thalamic domain, but the intracortical projections to the same neurons differed dramatically, with those to layer 6b originating from different laminar regions than those to layer 4.

Conclusion
Our results suggest that the intracortical projections to layer 6b likely contribute more to their complex receptive fields, while the thalamocortical inputs to layer 6b instead may be concomitantly attenuated. As such, the functional circuitry of the thalamocortical network is comprised of concurrent and convergent networks that lead to computationally divergent outcomes emerging from the intracortical network.
Subcollicular sources of projections to the auditory thalamus in the rat
Albert S. Berrebi1, F.-Javier Bernardo2, Yongming Jin3, Marcelo Gomez-Alvarezm, M-Auxiliadora Aparicio2, Emmanuel Marquez2, David Sloan1, Enrique Saldana2,4
1West Virginia University, 2University of Salamanca, 3Cleveland Clinic, 4Institute of Biomedical Research of Salamanca (IBSAL)

Background
It is generally assumed that the main source of ascending projections to the auditory thalamus is the inferior colliculus (IC). However, several isolated reports suggest that the medial geniculate body (MGB) receives direct projections from different auditory subcollicular nuclei (e.g. Henkel, Brain Res 259:21–30, 1983 [cat]; Angelucci et al., J Comp Neurol 400:417–439, 1998 [ferret]; Malmierca et al., J Neurosci 22:10891–10897, 2002 [rat]; Berrebi et al., ARO Abstr 35:250, 2012 [rat]). This is particularly relevant because evidence presented at this meeting indicates that, contrary to the long-held tenet, the medial division of the MGB (MGBm) is not a major target of the IC (Saldaña, Symposium on "The non-lemniscal auditory thalamus").

Methods
To systematically investigate the sources of ascending projections to the auditory thalamus, we made iontophoretic injections of the sensitive retrograde tracer FluoroGold into the auditory thalamus and analyzed the distribution of the neuronal cells bodies labeled in all subcollicular auditory nuclei.

Results
We obtained cases with injection sites centered in different regions of the auditory thalamus, including the ventral, dorsal and medial divisions of the MGB, the posterior intralaminar nucleus, the marginal zone and the posterior limitans nucleus. In the classical auditory nuclei, abundant retrogradely labeled neurons were found in the contralateral dorsal cochlear nucleus and lateral superior olive; in the ipsilateral medial superior olive (MSO), superior paraolivary nucleus (SPON), and ventral nucleus of the lateral lemniscus; and in the dorsal nucleus of the lateral lemniscus and the nucleus sagulum of both sides. We also found numerous labeled neurons ipsilaterally in the ventrolateral tegmental area, located medial to the ventromedial portion of the VNLL, as well as in an ill-defined territory wedged between the medial nucleus of the trapezoid body and the SPON. In the cases with the most retrograde labeling, the percentage of labeled neurons in some nuclei, such as the ipsilateral MSO and SPON, approached 50%.

Conclusion
Because FluoroGold injection sites tend to be large and usually include more than one subdivision of the auditory thalamus, we are currently performing a quantitative, multivariate analysis to infer the relative contribution of each one of the subcollicular nuclei to the innervation of each region of the auditory thalamus.

These findings improve our understanding of the organization of the auditory pathway, shed light on the complex parcellation of the auditory thalamus, and provide novel morphological frameworks for future functional studies.

Modulation of auditory steady-state response (ASSR) as a function of frequency ratio characterizing three-note chords.
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Background
Chords composed of tones characterized by simple frequency ratios are perceived consonant whereas chords of complex ratios are perceived dissonant. In theory, perception of dissonance is attributed to sensation of beats/roughness. The neuronal processing of beats/roughness is reflected in the auditory cortical steady-state response (ASSR), an oscillatory evoked response phase-locked to the temporal envelope periodically amplitude-modulated at rates of difference frequencies (delta-F). Intracranial recording revealed that the ASSR was correlated to energy of delta-F and exhibited larger activities for dissonant than consonant chords. However, effect of frequency ratio on the ASSR was yet unclear because the physical parameters of delta-f contained in each chord differed in terms of modulation rate, number of frequency components and spectral energy. The present study therefore aimed at exploring more simply the modulation of neuromagnetic ASSR as a function of frequency ratio.

Methods
Six kinds of stimulus chords composed of three sinusoids were prepared. The frequency ratio was varied systematically from 4:5:6, 6:7:8, through to 14:15:16, corresponding to chords containing 5th, 7th, to 15th overtone series. The carrier
frequencies of each chord were defined for the envelop to be amplitude-modulated at delta-F of 40 Hz. The spectral energy of all frequency components was equalized at 70 dB SPL by real-ear measurement and inverse filtering method. A sinusoidally amplitude-modulated (SAM) chirp tone was presented as a control to capture the neuronal frequency characteristics of the ASSR.

**Results**
Clearly elicited 40-Hz component was identified as the ASSR. Equivalent current dipole (ECD) of the ASSR was estimated using a single dipole model per hemisphere. The strength of the ECD moment for 5th and 7th chords characterized by simple frequency ratio was significantly smaller than that for 11th, 13th and 15th chords of complex frequency ratio despite that the acoustic parameters, i.e., the frequency as well as the spectral energy of delta-F to which the ASSR was synchronized was physically equal and further that the neuronal frequency characteristics was compensated by responses to the SAM chirp tone.

**Conclusion**
The results of the present study suggest that information that exists in natural world such as simple frequency ratio is processed in energy-saving manner whereas artificial information induces larger activations. Attentional gating of thalamocortical function as well as resonance effect with the spontaneous rhythmic activities might be candidates for the underlying mechanism.

MANTA – An Open-Source, High Density Electrophysiology Recording Suite for MATLAB
Bernhard Englitz$^{1,2}$, Stephen David$^3$, Shihab Shamma$^{1,2}$
$^1$Ecole NormaleSuperieure, $^2$University of Maryland, $^3$OHSU

**Background**
The distributed nature of nervous systems necessitates to record from a large number of sites in order to break the neural code, whether single cell, local field potential (LFP), μECOG, electroencephalographic (EEG), magnetoencephalographic (MEG) or in vitro micro-electrode array (MEA) recordings are considered. High channel-count recordings also optimize the yield of a preparation and minimize the time-investment for the scientist. Currently, data acquisition (DAQ) systems with high channel counts (>100) can be purchased from a limited number of companies at considerable prices.

**Methods**
We have developed an open-source MATLAB-based DAQ system MANTA (MAtlab N-Times Analog) which combines high channel counts (up to 1500 channels/PC), usage of analog or digital headstages, low per channel cost (<$90/channel), feature-rich display & filtering, a user-friendly interface, and a modular design permitting easy the addition of new features. MANTA is licensed under the GPL and free of charge.

**Results**
The system has been tested by daily use in multiple setups for >1 year, recording reliably from 128 channels. It offers a growing list of features, including integrated spike sorting, PSTH and CSD display and fully customizable electrode array geometry (including 3D arrays), some of which are not available in commercial systems. MANTA runs on a typical PC and communicates via TCP/IP and can thus be easily integrated with existing stimulus generation/control systems in a lab at a fraction of the cost of commercial systems.

**Conclusion**
With modern neuroscience developing rapidly, MANTA provides a flexible platform that can be rapidly adapted to the needs of new analyses and questions. Being open-source, the development of MANTA can outpace commercial solutions in functionality, while maintaining a low price-point.
Cytoarchitecture and Parvalbumin Immunoreactivity of the Auditory Cortex in the Chinchilla.

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Background
Although chinchillas (Chinchilla laniger) have been widely used as a model to study middle ear and cochlear anatomy, there are relatively few studies focused on auditory cortex morphology. The bifurcation of the middle temporal artery has been proposed as a vascular landmark to identify primary auditory cortex in chinchillas. In addition neuronal responses to brief tones with a latency <15 ms, have been postulated to be generated in the primary auditory cortex of chinchillas. Here, we measured auditory cortex evoked potentials and correlate these findings with cytoarchitecture (Nissl) and parvalbumin immunoreactivity of the auditory cortex.

Methods
Ten adult chinchillas were anesthetized and placed in a stereotaxic frame inside a sound attenuated room. Macroscopic vascular landmarks were measured from bregma. An electrophysiological characterization of the auditory cortex was performed using tones at different frequencies (1-8 kHz) and intensity levels (20-80 dB SPL). An electrolytic cortical lesion was made by a current pulse (1 mA for 15 s) throughout the recording electrode and the cytoarchitecture (Nissl) and immunohistochemistry with Parvalbumin were evaluated.

Results
The bifurcation of the middle temporal artery was found in 8 animals, located in average at: 2.6 ± 0.9 mm (X axis), 10.0 ± 1.9 mm (Y axis), and 3.9 ± 1.3 mm (Z axis) as measured from bregma. The response latency of this brain position was shorter than 15 ms only in three animals. The average cortical thickness of sites with latencies < 15 ms was 2070 ± 119 µm, while that of sites with latencies between 15-20 ms was 2230 ± 279 µm (p>0.05). Both auditory cortices were thicker than parietal sensory cortex (1395 ± 187 µm; p<0.001). These thickness differences were mainly due to thicker layers V and VI in both auditory fields (p <0.01). The density of parvalbumin (+) neurons was similar in parietal and auditory cortex exhibiting higher counts in layers IV and V.

Conclusion
The primary auditory cortex of the chinchilla was located in the bifurcation of the middle temporal artery in 30% of the experiments. A noticeable feature of the chinchilla auditory cortex was its thickness around 2 mm, which depends mainly on thick layers V and VI. Parvalbumin (+) immunoreactivity was found mostly in layers IV and V. To guarantee the exact location of the primary auditory cortex in chinchillas, electrophysiological confirmation is needed in every experiment.

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Middle Latency Auditory Evoked Potentials in Response to Feature-Conjunction Deviants suggest Processing Downstream to Simple Deviant Processing

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Background
The violation of a regular sound pattern by irregular or novel stimuli is reflected by the mismatch negativity (MMN), a component of the human auditory evoked potential (AEP), with a latency of 100-250 ms and main sources in the auditory cortices. Evidence for auditory novelty detection however has also been reported in the middle-latency range (MLR) of the human AEP at latencies from 20 to 40 ms after change onset, in particular for simple feature deviants. According to a hierarchical view of auditory novelty detection, more complex types of regularities should be encoded at higher levels of the auditory pathway than simple regularities. In fact, MMN can be triggered by a so-called feature conjunction deviant, which combines features of different standard stimuli, e.g. the frequency of one standard and the location of another standard. In this study, we aimed at examining if the detection of conjunction deviants becomes already evident in the MLR of the human AEP.

Methods
A conjunction paradigm was applied (standard 1 [45%] = 800 Hz, presented from the left side; standard 2 [45%] = 1200 Hz, presented from the right side; deviant 1 [5%] = 800 Hz, presented from the right side; deviant 2 [5%] = 1200 Hz, presented from the left side), while subjects were concentrating on a subtitled movie. In a second session a frequency oddball paradigm was presented, in order to compare the deviance-related modulations of the AEP in response to a simple and a more complex regularity violation. The electroencephalogram was recorded from 64 electrodes. Different frequency filters for the analysis of the MLR and the long-latency range AEP were applied. The data of 17 subjects was statistically tested for differences in amplitudes and latencies of the MLR components and MMN elicited by standard and deviant stimuli.

Results
A significant MMN was obtained at fronto-central electrodes for both experimental paradigms. MMN in the oddball condition was significantly bigger than MMN in the conjunction condition. In the middle-latency range, the components Na, Pa and Nb showed no deviant-related modulations in response to the conjunction deviant. In the oddball condition, however, the amplitude of the Nb component was enhanced in response to the deviant, compared to the standard.

Conclusion
This outcome suggests that the detection processes of an auditory feature conjunction deviant start at auditory cortical areas lying downstream to the areas processing a simple auditory deviant.

Early deviance detection within an intensity pattern regularity: an electrophysiological study in humans

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Background
Recent oddball studies showed that auditory change detection responses exist in the first 50 ms after sound onset, in the middle latency response (MLR) range. These deviance-related MLR responses occur much earlier than the Mismatch negativity (MMN), an electrophysiological correlate of the brain’s pre-attentive auditory change detection system, observed as an auditory evoked potential at 100-250ms. However, when complex paradigms were employed, these early deviance effects were not as consistent, suggesting that different levels of information were encoded in two separate time ranges, hence supporting the notion of a functional hierarchy of auditory deviance detection. In the present study, we further tested this notion by examining if such early responses could be elicited by changes within an intensity pattern.

Methods
In order to obtain optimal MLR, we used an “up-chirp” stimulus that was designed to compensate for the cochlear traveling wave delay. It starts with a low frequency (50Hz) and ends with a high frequency (8000Hz), with a duration of 16.7ms. Electroencephalography (EEG) data were obtained from sixteen healthy participants. Participants were presented with a sequence of chirps that followed a rising intensity pattern (55, 65, 75, 85dB). This pattern was violated 15% of the time, with a pattern in which the third and the fourth tones had the same intensity (55, 65, 75, 75dB). We
compared the responses elicited by the third (standard) and the fourth (deviant) chirps within the rare pattern, in both MLR and MMN latency ranges.

**Results**
In the MLR range, the deviant led to an enhanced Pa response (corrected p = 0.030) at around 42 ms when compared to the standard. No significant differences were observed between deviant and standard responses for other MLR components (Na, Nb). In the MMN range, we observed a double-peaked negativity on the difference waveform (deviant minus standard) at 78-108ms and at 175-205ms respectively.

**Conclusion**
Our results suggest that the human brain is capable of detecting an auditory pattern change as early as 42ms. Although there is a possibility that the observed effect could be due to repetition enhancement, it is unlikely to be the case, especially since repeated stimuli have been shown to reduce neuronal responses, as demonstrated by stimulus-specific adaptation in animals and sensory gating in humans. The present study has shown, for the first time, that deviance detection within an intensity pattern regularity could be reflected electrophysiologically in human MLR.

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**Neuromechanistic Temporal Models of Pitch Perception Using Slope Detectors**
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**Background**
Despite the importance of pitch in auditory scene analysis, the neural mechanisms for pitch detection remain unclear. The existence of pitch-selective neurons in nonprimary auditory cortex suggests a separate pathway for pitch processing and a possible pitch map in cortex (Bendor & Wang 2005). Most existing pitch models are one of two types: pattern matching (spectral) or autocorrelation (temporal). While these models can predict pitch in many cases, development of neuronal implementations is immature. We have developed two mechanistic temporal models using two different slope detector units, extracting pitch information explicitly as a place code and temporal spike trains respectively.

**Methods**
We developed a firing rate (mean field) model and a biophysical cellular (HH-like) model each with phasic behavior, i.e. each has an onset-only response followed by low steady state during the stimulus regardless of stimulus amplitude (Figure1). These models can serve as slope detectors, since stimulus needs to rise fast enough to initiate a response. The population model has a graded response and resonant frequency, while the cellular model can produce precise phase-locking. Using a tonotopic array of such (uncoupled) units with a broad footprint for afferent input, the models can respond at the period corresponding to the perceived pitch as reported in human psychoacoustic studies.

**Results**
Our models can encode the fundamental frequency of missing fundamental harmonic complexes while being unresponsive to individual components when presented alone. The response amplitude decreases as the lowest order of harmonics increases, consistent with the decrease in salience of human pitch perception. For shifted harmonics, our models also reproduced the perceived pitch which is not the fundamental frequency. Moreover, our model is not sensitive to phases of the harmonics. With alternating phase and Schroeder phase our results are consistent with psychoacoustic studies (Figure2). Our population models represent the extracted pitch in two ways. The resonant units have preferred pitch, thus representing pitch with a place code. The phase-locking units encode pitch as temporal periodic firing, in which case another stage is needed to transform into rate code.

**Conclusion**
Using an array of slope-detector units that receive input from broad frequency channels, our models can extract pitch not only for missing fundamental complexes, but also encode correctly in special cases when pitch does not correspond to periodicity or envelope. Different from the pattern matching and autocorrelation models, our models do not require a bank of coincidence detectors and delay lines.
Combinatorial regulation of hair cell induction by co-transfection of ATOH1 and transcription factors.
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Background
Overexpression of the basic helix-loop-helix transcription factor (TF) Atoh1 has been shown to induce the transformation of supporting cells in the organ of Corti into hair cells (HCs). Evaluating potential regulatory region of the pou4f3 gene, a likely target of Atoh1 in HCs, we identified a cluster of binding sites for Atoh1 and a number of other TFs, a feature that was highly conserved across a widely separated mammalian species. We previously tested three of these (TCF3, GATA3 and SP1) and found that two (TCF3 and GATA3) enhanced the ability of Atoh1 to activate the gene and induce Myo7A expression in nonsensory cells.

Methods
To test the hypothesis that the remaining TFs act in combination to regulate the pou4f3 gene, we transfected by electroporation neonatal mouse (postnatal day 1.5) organ of Corti with plasmids encoding these TFs for which highly conserved binding sites have been identified in the pou4f3 gene and which are expressed in the developing organ of Corti: CUX1, E2F1, ETS2, ETV4, FOXM1, HES1, HES5, GABPA, GATA1, GATA2, MAX, MAZ, MEIS, NERF, NFE2, NMYC, PATZ1 and USF2. We used a transgenic mouse in which 5′ f DNA from the pou4f3 ATG drives expression of GFP to assess the regulation of this gene, and labeled the cultures with anti-myosin 7A to identify transition to a HC-like phenotype.

Results
We identified several additional TFs had enforced the effect of Atoh1 while others inhibited or did not influence Atoh1.

Conclusion
These results suggest that Atoh1 acts in concert with a subset of other TFs that find that find to adjacent DNA element to control the expression of the pou4f3 gene, and more generally to induce a HC phenotype. These TFs could be used to enhance HC regeneration.

Tonotopy organization of the vertebrate cochlea may require a gradient of Shh signaling
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Background
Precise perception of sounds is crucial for verbal communication as well as daily activities of human life. Frequency discrimination first occurs at the peripheral auditory organ, the cochlea, which is tonotopically organized such that hair
cells are tuned to high frequency sounds at the base of the cochlea and to low frequencies at the apex. However, the underlying mechanisms for tonotopy remain obscure. It has been shown that graded levels of Sonic hedgehog (Shh) signaling provide unique positional information in various tissues including neural tube and limbs. Since the developing cochlea receives graded Shh signaling along the cochlear duct, higher at the apex and lower at the base, it has been suggested that the gradient of Shh signaling may confer unique positional identities along the cochlear duct, contributing to the establishment of tonotopy.

Methods
To test this hypothesis, we upregulated Shh signaling in the developing mouse cochlea using the cre/lox approach as well as implanting Shh-soaked beads to the developing chicken cochlea.

Results
When Shh signaling was ectopically activated in the cochlear duct of Pax2<sup>Cre/+</sup>; Smo<sup>M2/+</sup> mutants, genes that are preferentially expressed in the apical cochlear regions such as Fst, Msx1, and EphrinB2, were upregulated throughout the cochlear duct, whereas expression of A2m, which is normally restricted to the base, was downregulated. These results suggest that Shh may specify regional identity of the developing cochlea. Hair cell phenotypes cannot be evaluated in these mutants due to early lethality. In contrast, implanting beads soaked with Shh proteins into the chicken otocyst <i>in ovo</i> generated hair cells at the base of the basilar papilla with stereocilia morphologies that resembled those located more apically in controls.

Conclusion
Our results suggest that high Shh signaling promotes apical cochlear identity and a gradient of Shh signaling may facilitate the establishment of the tonotopic organization of the auditory organ in vertebrates.
PTEN plays multiple roles by regulating both PI3K/Akt and MEK/Erk signaling pathway during the mammalian inner ear development

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Background
The PI3K/Akt signaling pathway has been shown to integrate multiple developmental signals to promote cell growth, proliferation and survival, and these effects of PI3K/Akt signaling are counteracted by the tumor suppressor PTEN (phosphatase tensin homologue) through its phosphatase activity. In this study, we investigate the regulatory role of PTEN in mammalian inner ear morphogenesis, using mice in which PTEN gene is specifically eliminated from developing inner ear.

Methods
To disrupt PTEN function within the inner ear, we crossed PTEN<sup>flox</sup>/PTEN<sup>flox</sup> mice (Suzuki A, Mak TW et al., 2003) with doubly heterozygous mice for the Foxg1-Cre allele (Hebert JM, McConnell SK, 2000) and the PTEN-<sup>flox</sup> allele. Offspring with the genotype Foxg1<sup>Cre</sup>/+; PTEN<sup>flox</sup>/PTEN<sup>flox</sup> (PTEN-cko) survived through E18.5, and the immunoreactivity of PTEN had disappeared and the activation of Akt was indeed increased in PTEN conditional knock-out (cko) inner ear.

Results
PTEN cko inner ear was hypertrophic, and especially, the diameter of cochlear duct had extraordinarily increased. The increased number of PHH-positive, proliferating cells implicated that the cell proliferation was prolonged in PTEN cko inner ear. In PTEN cko littermates, the extra OHCs are found in patches that alternated with regions containing the normal three rows. The additional IHCs appear in places, however, less frequently compared to the case of OHCs. As Myosin VI-immunopositive OHCs were found only in the apical-middle turn of PTEN cko cochlea and not in that of wild type cochlea, the differentiation of HCs seemed to be accelerated in PTEN cko mutants. The extent and relative positioning of Sox2 and p27Kp1 expression were unchanged in PTEN cko mutants at E13.5, which imply that the prosensory formation is almost normal in PTEN cko cochlea. The expression of activated Akt was increased by deletion of PTEN, especially in the neurons and the out side of the prosensory region. Recent reports have showed that PTEN also negatively regulates MEK/Erk pathway. Indeed, activated Erk was expressed in the supporting cell progenitors at E15.5 and E17.5 in PTEN cko mutants, which was not observed in WT controls.

Conclusion
The multiple roles PTEN were clarified by analyzing the inner ear of PTEN cko mouse. PTEN regulate the MEK/Erk signaling pathway, in addition to the PI3K/Akt signaling pathway. PTEN controls the premature differentiation of HCs and the prolonged continuous cell proliferation to complete the adequate mammalian inner ear morphogenesis.
Characterization of Hippo Signaling During Development and Regeneration of Inner Ear Sensory Hair Cells
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Background
The loss of sensory hair cells (HCs) is a leading cause of deafness in humans. Although the mammalian inner ear is unable to replace lost/damaged HCs, birds and other non-mammalian vertebrates can regenerate HCs after injury. The Hippo signaling pathway constitutes an evolutionarily conserved kinase cascade involved in regulation of organ size, cell proliferation, and tissue regeneration. Despite these well-documented roles in growth control, Hippo signaling remains unexplored in the ear. The purpose of this study is to begin characterization of the Hippo pathway in the context of inner ear biology, with focus on development and regeneration of sensory HCs.

Methods
Immunocytochemical techniques were used to characterize the expression patterns of several core Hippo signaling components (e.g. SAV1, NF2/Merlin, MST1/2, LATS1/2, YAP1, and TAZ) within the developing mouse inner ear. Illumina sequencing was used to identify expression of Hippo pathway genes in organotypic cultures of normal and Streptomycin-damaged chick utricles and cochleae.

Results
Temporal changes in the distribution of several Hippo signaling components, including SAV1, NF2/Merlin, and YAP1 indicate that the Hippo pathway may be involved in the regulation of terminal mitosis and/or onset of HC differentiation during mouse organ of Corti development. In mature avian sensory epithelia, RNA sequencing revealed robust expression of numerous canonical Hippo signaling components, including SAV1, STK3/4, LATS1/2, NF2, YAP1, TAZ/WWTR1, DIAPH1/3, DLG2, and MOB2, as well as the upstream Hippo signaling regulators alpha-catenin, FAT1-4, and DCHS1/2. Several TEAD family transcription factors (1, 3, and 4), which are required for YAP1-mediated induction of proliferation, demonstrated significant expression levels as well. Finally, sequencing data revealed alterations in YAP1 target gene expression (e.g. MYC, DIAPH1, TP53BP2, TP63, TP73, E2F1, and Cyclin E) throughout the time course of avian HC regeneration in vitro.

Conclusion
Despite its roles in cell proliferation and tissue regeneration in other organ systems, Hippo signaling has not been studied in the inner ear. The data presented here describe the localized expression of key Hippo signaling molecules within sensory epithelia of avian and murine inner ears, and provide evidence to support putative roles for Hippo signaling during sensory epithelial development and HC regeneration.

Age-related changes in the expression of vesicular glutamate transporter 3 in the rat cochlea
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Background
Vesicular glutamate transporters 3 (VGLUT3) plays an important role in hearing maintaining, and knockout mice in which are totally deaf and auditory brainstem responses (ABRs) indicates an early defect. Yet the exact mechanism in the cochlea is not well established. Although the presence of VGLUT3 in fetal and postnatal auditory hair cells are confirmed, the age associated changes of VGLUT3 expression in aging animals remains unclear and the research doesn’t involve spiral ganglion cells (SGCs). Elucidation of developmental and aging dynamics of VGLUT3 expression in the auditory epithelium and SGCs is conducive to functionally understanding auditory glutamatergic transmission.

Methods
We characterized the expression profile of VGLUT3 in the rat cochlea and SGCs during postnatal development and aging by using immunohistochemistry and quantitative real-time polymerase chain reaction (qRTPCR).

Results
We found that the expression of VGLUT3 in the cochlea varied from different age groups. VGLUT3 expressed in the inner hair cells (IHCs) and SGCs in the cochlea as early as the post-natal days, and present a dynamic variation during the development. We noted that postnatal 14 days (P14) was a key time point when the expression of VGLUT3 in IHCs sharply increased and VGLUT3 expression peaked in the adult. VGLUT3-immunopositive puncta was found in the cytoplasm of SGCs from P7 but not in IHCs. And VGLUT3 was also detected in the cochlea of aging rat. In IHCs, no
apparent difference of VGLUT3 expression was found between the aging and adult rat. Although in SGCs, the amount of SGCs decreased in comparison with that of adult, the fluorescence of VGLUT3 in SGCs present little change with the method of immunohistochemistry. Slc17a8 gene expression trend evaluated by qRTPCR is similar with that of the VGLUT3 protein expression. And no statistic difference was found in the mRNA expression level between the groups of adult and aging rats.

Conclusion
VGLUT3 expression of the cochlea exists age-related changes. At the hearing onset stage (P14), its expression in IHCs is obviously intense and reach peak at the adult. VGLUT3-immunopositive puncta was found in the cytoplasm of SGCs from P7 but not in IHCs. Except for the decreasing with the SGCs loss, there are no obvious VGLUT3 expression changes in the aging rat cochlea. Since it is present in early developmental stages and its expression increases with maturing and still maintains to the aging, VGLUT3 is likely to have developmental and physiological roles in rat cochlear.
Characterization of Myosin7a in the Cochlear and Vestibular Ganglia of the Developing Chick
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Background
The cochlear vestibular ganglion (CVG) comprises a cluster of undifferentiated neurons that delaminate beginning around embryonic day 2 (E2). The CVG will then divide into two separate populations; the cochlear and vestibular ganglion by (E6.5). Various transcription factors have been shown to play a role in the differentiation of both cochlear and vestibular neurons in mice (Kim et al., 2001; Lawoko-Kerali et al., 2004; Lu et al., 2011). Previous work from our lab has demonstrated a possible role for NeuroD and Gata3 in influencing neuroblast cell fate decision in chicks (Jones and Warchol, 2009).

Methods
We utilized immunohistochemistry to characterize the expression of Myosin7a in the developing chicken inner ear neurons from E3 to E11.

Results
We are investigating whether Myosin7a, an actin-dependent molecular motor protein, may play a role in the decision of a neuronal precursor to become a cochlear or vestibular neuron. The onset of Myosin7a expression was detected at E4 in a subset of neuronal precursors. Expression becomes restricted to vestibular neurons at E5 and persists as late as E7.

Conclusion
Our results indicate that Myosin7a may be playing a role in the development of vestibular neurons.
Septin7 Expression and Function in the Inner Ear of Embryonic Mouse
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Background
Septin proteins are GTP-binding proteins that are evolutionally conserved through eukaryotes except plants. Septin complex can interact with membrane, myosin, and microtubules. Their functions include formation of the cortical rigidity, control of the vesicular transportation, and compartmentalization within plasma membrane. These functions contribute to the formation of the cellular polarity that is important characteristics of epithelial cells. Our previous reports showed that Septin7, a core protein of multimeric Septin complex, was expressed in the embryonic cochlea (Yoshida et al. Hear Res 2012).

Methods
To identify the roles of Septin7 in the development of inner ears, we crossed Septin7 floxed mice with Foxg1Cre mice because conventional Septin7 knockout mice were embryonic lethal. Then we investigated the inner ear morphology of the mice using paint filling and the expression patterns of several markers related to early inner ear development.

Results
Macroscopic morphology by paintfilling on E17.5 inner ears showed remarkable underdeveloped inner ear with only cystic structure. Normal otocyst was formed in E10.5 but later development of the inner ear goes on abnormally.

Conclusion
This result shows that Septin7 plays significant role in early developmental stage of inner ear.
Exploring MicroRNA Expression Patterns in Zebrafish and Chicken Inner Ear
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Background
MicroRNAs (miRNA) are small single-stranded RNAs that mediate post-transcriptional repression of gene expression by binding with target mRNA 3'UTRs. Mice with conditional knockouts of Dicer1 in embryonic inner ear have severe neurosensory defects and hair cell degeneration in three studies, revealing the importance of miRNAs in hair cell maturation and maintenance. Using published data from miRNA expression screens in mice, as well as miRNA target gene prediction algorithms, we created a list of miRNAs to interrogate for cross-species conservation of expression in the inner ear. Our aim was to use these model organisms to understand the roles of individual miRNAs in the developing inner ear.

Methods
Digoxygenin-labeled locked-nucleic acid probes for miR-19b, 93, 99, 124, 130b, 141, 199, 200a, 200b and 200c were purchased from Exiqon. We performed whole-mount in situ hybridization of zebrafish embryos at 1, 2, 3 and 5 days post fertilization (dpf). A similar method was applied to whole mounts of the chicken basilar papilla (BP) on embryonic day 17 to localize miR-96.

Results
We found that miR-200a and miR-141 were weakly expressed in inner ear hair cells and highly expressed in lateral line neuromasts and olfactory epithelia. At 2dpf and 5dpf, miR-124 gave strong signal in the eye. Broad expression in the head was observed for miR-93. We were unable to detect expression of miR-19b, 99, 130b or 199 in zebrafish at the tested ages. In the chicken BP, miR-96 was expressed in hair cells with a pronounced longitudinal gradient (high apex, low base).

Conclusion
We confirmed the evolutionary conservation of inner ear expression for the miR-200a family in zebrafish, adding miR-141 to the list of genes associated with mechanosensory organs. However, it is proving challenging to identify additional miRNA families associated with the inner ear in zebrafish. In chicken embryos, the miR-96 expression gradient (apex higher than base) matched that reported for the mouse cochlea after 2 weeks of age (Weston et al., 2011). In mice, this pattern for the miR-183 family emerged from an inverse gradient at postnatal day 0 (base higher than apex). A developmental timecourse of expression for the BP is underway to ask if this spatiotemporal expression pattern is also conserved. Expression analysis of miR-9 in the BP is also ongoing.

Temporal-Spatial Expression of microRNAs in Mouse Inner Ear Sensory Epithelium
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Background
MicroRNAs (miRNAs) post-transcriptionally regulate gene expression and are required in the development of the inner ear. Previous studies demonstrate that ~100 miRNAs are expressed in mouse inner ear. Certain of these miRNAs exhibit hair cell specific expression including the miR-183 family (miR-183, miR-96, and miR-182) that is known to impact hair cell differentiation and maintenance. However, little is known about the remainder of inner ear miRNAs. To gain greater perspective on cell-specific expression and function of inner ear miRNAs, we have completed an expression survey of 76 miRNAs.

Methods
In situ hybridization using a digoxigenin (DIG)-labeled, locked nucleic acid (LNA) probe complementary to each miRNA was performed on embryonic or neonatal mouse inner ear. miRNA expression was visualized using anti-DIG antibody conjugated to alkaline phosphatase and BM purple as a chromogenic substrate. Tissues were whole-mounted or sectioned for microscopic imaging.

Results
Many miRNAs are detected in the mouse inner ear by microarray analysis and quantitative PCR. Our survey of miRNA expression patterns using in situ hybridization on neonatal mouse inner ear showed that many such miRNAs were ubiquitous with high or low detection, or undetected. Relatively fewer miRNAs or miRNA families showed cell-specific expression patterns that included sensory neurons or epithelia. Of particular interest are those miRNAs that exhibited...
sensory epithelium and hair cell-specific expression in embryonic and neonate mouse inner ear. These included miR-9, miR-200 family (miR-141, miR-200a, miR-200b, miR-200c, miR-429), and miR-204 family (miR-204, miR-211) in addition to miR-183 family.

**Conclusion**
Results suggest that relatively few yet highly conserved miRNA families contribute exclusively to the development of sensory epithelia and hair cells in the mouse inner ear, whereas many other miRNAs might serve relatively general albeit important functions.

**Keratan sulfate mediates zebrafish otolith formation**  
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**Background**
More than 50 human proteins have been identified in which mutations are associated with congenital hearing loss. While many of these proteins have glycans attached to them, little is known about the role these sugars have in mechanosensory function. The glycosaminoglycan keratan sulfate (KS) is consistently found in the extracellular matrixes that directly overlie sensory hair cells and are critical to mechanosensory function. We sought to define the expression of KS during early zebrafish otolith development, identify which enzymes were responsible for KS synthesis, and characterize otolith development in KS-deficient zebrafish embryos.

**Methods**
Zebrafish embryos were collected for KS-immunohistochemistry at developmental stages when otic structures were first visible, including the otic placode (16 hours post fertilization, hpf), otocyst (18 hpf), and otolith (19.5 hpf). Early stages of otolith development were also examined at 21, 24, and 27 hpf. *In situ* hybridization with probes for putative KS-synthesizing enzymes was done using embryos collected at the same stages. Morpholinos were injected at 1-cell stage, and otoliths were examined at 27 hpf.

**Results**
KS is expressed at the earliest stages of zebrafish otolith development. While KS is actively synthesized by the cells surrounding the otocyst beginning at 16 hpf, KS is not found in the otolith until 24 hpf, approximately 6 hours after the earliest otolith-localized protein, Oc90. KS forms a layered biomatrix that complements the expression of the otolith matrix protein OMP1. We identified *b3gnt7b*, *chst2b*, and *chst1* as putative KS-synthesizing enzymes there were expressed in the otic placode concomitant with early KS expression. Morpholino knockdown of either *b3gnt7b*, *chst2b*, or *chst1* resulted in embryos with deformed or absent otoliths.

**Conclusion**
During early zebrafish otolith development, KS synthesis and secretion are temporally distinct. We identified genes responsible for three of the four steps of KS synthesis during zebrafish otolith development; and using morpholino knockdown of each gene, we have demonstrated that KS is required for zebrafish otolith formation.
The Role of the Notch Signaling Pathway in Supporting Cell Development

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Background

The sensory epithelium of the cochlea is responsible for the perception of sound. This specialized epithelium consists of sensory hair cells and glial-like supporting cells. Auditory supporting cells provide structural and functional support to hair cells and are critical for cochlea homeostasis and function. Despite the fact that these cells are functionally important to hearing and a possible unit of regeneration within the inner ear, little is understood about the molecular mechanisms involved in their differentiation. A good candidate pathway for being involved in supporting cell differentiation is the evolutionarily conserved Notch signaling pathway, which is highly activated in the sensory progenitors fated to differentiate into supporting cells. It is well established that activation of the Notch signaling pathway in these sensory progenitors results in the up-regulation of Atoh1 antagonists preventing these progenitor from adopting a hair cell fate. We hypothesize that the Notch pathway, in addition to repressing the hair cell fate in bi-potential progenitor cells, promotes supporting cell differentiation by up-regulating genes important for this process.

Methods

We used DAPT, a gamma secretase inhibitor, to disrupt the Notch signaling pathway in cultured developing cochlea (E15.5) for 18 hours. We analyzed the resulting changes in gene expression with Affymetrix exon arrays and we used RT-qPCR for microarray data validation. Using RT-qPCR we analyzed supporting cell specific gene expression in the Cre inducible dominant negative mastermind like 1 (DN-MAML1) transgenic mice. The expression of DN-MAML1 disrupts Notch signaling by preventing the formation of the activated Notch transcriptional complex.

Results

Acute DAPT inhibition of the Notch signaling pathway uncovered more than 200 cochlear epithelial genes that are positively regulated by the Notch signaling pathway. Based on function and supporting cell specific expression we selected 24 genes for further validation in the DN-MAML1 transgenic mouse line. Our results indicate that a vast majority of the selected genes showed a decrease in expression in this Notch mutant.

Conclusion

In summary, our data shows that a significant subset of supporting cell specific genes are regulated by the Notch signaling pathway, suggesting that this pathway positively regulates supporting cell differentiation.

The Role of the Lin28b/Let-7 Axis in Cochlea Differentiation

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Background

The development of the mammalian auditory sensory epithelium is a carefully orchestrated process that results in the generation of several highly specialized cell types from a single progenitor cell pool. These mechanosensory hair cells and supporting cells are critical for hearing, but the molecular mechanisms regulating their differentiation are largely unknown. Using a high throughput screen we identified that the heterochronic gene Lin28b is selectively expressed in auditory sensory progenitors. This RNA-binding protein maintains cells in a pluripotent state by two distinct mechanisms: it enhances translation of growth-related genes and represses the expression of the let-7 family of microRNAs. We hypothesized that the Lin28b/let-7 miRNA axis is playing an important role in the transition of cochlear progenitor cells to a differentiated state.

Methods

We used both in situ hybridization and quantitative PCR to characterize the temporal and spatial expression pattern of Lin28b and the let-7 miRNAs during cochlea differentiation. In order to address the role of Lin28b in cochlea differentiation, we used both lentiviral-infected organotypic cochlea cultures and a doxycycline-inducible transgenic mouse line to overexpress Lin28b prior to hair cell differentiation.

Results

During cochlea development, Lin28b is expressed in the undifferentiated cochlea sensory progenitor cells, but drastically decreases as sensory progenitors differentiate. Conversely, let-7 miRNA expression increases with progressive sensory epithelium differentiation. Overexpression of Lin28b within the undifferentiated cochlea led to a significant slowing in the
timing of hair cell differentiation. While hair cells do eventually form in the Lin28b overexpressing cochlea, Lin28b overexpressing hair cells lag in their maturation and stereocilia formation.

Conclusion
We found that the Lin28b/Let-7 axis plays an important role in the timing of hair cell differentiation and maturation. Here we shall discuss the gene targets and molecular mechanisms underlying Lin28b’s role in cochlea differentiation.

Normal morphogenesis of auditory hair cells requires both non-muscle myosin IIA and IIB
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Background
In humans, the heavy chains of the non-muscle myosin II (NMII) family are encoded by three genes, MYH9 (NMIIA), MYH10 (NMIIB) and MYH14 (NMIIC). Mutations in either MYH9 or MYH14 cause hearing defects. The mechanisms of these defects are unclear. NMII has been shown to play significant roles in junction formation and cell shape changes. Recent studies have suggested that NMII is involved in cochlear duct extension and patterning during the organ of Corti development. However, it is still not clear what roles NMIIA and NMIIB have during hair cell development and maturation.

Methods
To investigate the roles of NMIIA and NMIIB in hair cells, we used the Cre-loxp system to specifically delete (cKO) NMIIA and NMIIB in hair cells since both NMIIA and NMIIB germline knockout mice are embryonically lethal. We bred NMIIA floxed and NMIIB floxed mice with the hair cell specific Cre mouse line Gfi1-Cre to generate NMIIA cKO, NMIIB cKO and NMIIA and NMIIB double cKO mice. Auditory brainstem responses (ABR) were monitored in each mouse model. Whole mount immunohistochemistry was performed for hair cell morphologic analysis.

Results
Neither NMIIA cKO nor NMIIB cKO mice showed any ABR hearing defects or detectable morphological defects in the organ of Corti. NMIIA and NMIIB double cKO mice show significant hearing defects and abnormal hair bundle shapes. These abnormal hair bundles do not have the normal V-shape, instead they show an asymmetric V-shape or S shape.

Conclusion
Normal morphogenesis of auditory hair cells requires both non-muscle myosin IIA and IIB.

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Plastin-1 Controls Vestibular Stereocilia Length

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Background
The mechanosensory hair bundle consists of organized rows of actin-filled stereocilia of increasing heights. During development, stereocilia form through the assembly of actin filaments into tight parallel bundles by actin-bundling proteins. Stereocilia then increase in diameter and length through the addition of new actin filaments and the elongation of existing filaments within the actin paracrystal. Although plastin-1 (PLS1) is a member of one of the three classes of actin-bundling proteins known to be enriched in hair bundles, little is known about its function in stereocilia assembly.

Methods
We used mass spectrometry to identify and quantify proteins of hair bundles purified from P21-P25 mouse utricles. Protein abundance and localization was carried out with quantitative immunoblotting and confocal immunocytochemistry. Morphometric analysis of stereocilia was conducted with confocal images of phalloidin-labeled utricles.

Results
We tested the importance of PLS1 in stereocilia development by examining the vestibular hair bundles from Pls1-null mice. Pls1-null mice do not display behavioral evidence of overt auditory or vestibular dysfunction and auditory and vestibular hair cells appear to develop normally. To test whether loss of PLS1 is compensated by upregulation of another actin-bundling protein, we used mass spectrometry to quantify proteins in purified mouse hair bundles from wildtype and Pls1-null mice. In wildtype mice, PLS1 is the second-most abundant protein in the bundle and the most abundant actin-bundling protein of those we detected (PLS1, FSCN2 and ESPN). PLS1 was absent from the bundle in Pls1-null mice but no other actin-binding proteins increased in expression or were mislocalized. Using mass spectrometry and immunoblotting, however, we detected a significant ~30% decrease in the total amount of actin within the bundles of Pls1-null mice. Morphometric analyses of vestibular bundles indicated that stereocilia of adult (P21) Pls1-null mice were significantly shorter in length (~20%) than those of wildtype mice, with no change in kinocilia length or the staircase architecture of the bundle. Stereocilia at early postnatal stages (P4-P6) were also shorter than those of wildtype mice, although the difference was smaller (~10%).

Conclusion
Our results suggest that PLS1 is not essential for the initial assembly of stereocilia, but that it does play an important role in regulating the elongation of stereocilia. The difference in stereocilia between Pls1-null and wildtype mice was greater later in development, suggesting that PLS1 is used for stereocilia lengthening between P4 and P21.
Absence of Plastin1 results in a reduction of the width of inner hair cell stereocilia and a moderate hearing loss.

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Background

The stereocilia of hair cells contain a central core of F-actin filaments, organized into tightly packed parallel bundles. Proteins that bind to F-actin and promote such assembly are known as actin-bundling proteins, some of which such as espin, harmonin and eps8 are critical for stereocilia formation. However, the function in the inner ear of the most abundant actin-bundling protein expressed in stereocilia, plastin1 (pls1, also known as fimbrin), has not been examined. In the intestinal brush border of mice in which the gene encoding plastin1 has been disrupted (plastin1 KO mice) there are changes in microvillar dimensions, missing microvillar rootlets and reduction in the thickness and organization of the terminal web underlying the microvilli at the apical end of the cell (Grimm-Gunter et al. 2009). This suggests the possibility that stereocilia and auditory function may be affected in plastin1 KO mice.

Methods

The organs of Corti of plastin1 KO mice and their wildtype and heterozygous littermates (P8 to >6 months) were examined by immunohistochemistry, and scanning and transmission electron microscopy. Auditory function of adult mice (P30+) was tested by auditory brainstem responses to click and tone pip (8-40kHz) stimuli. Biophysical properties of hair bundle mechanotransduction were examined in organs of Corti from P6 animals.

Results

Hair cells developed normally in the vestibular and auditory epithelia of plastin1 KO mice. However, auditory brainstem responses revealed that plastin1 KO mice have a moderate hearing loss across most frequencies compared to their wildtype and heterozygous littermates. Ultrastructural studies of adult mice cochleae showed a significant reduction in the width of inner hair cell stereocilia but no effect on height, or general organization of the bundle. Outer hair cells showed no such defect. Cuticular plate depth and stereociliary rootlets of both hair cell types were also unaffected. In outer hair cells at P6, the extent of adaptation of mechanotransduction current was increased and fast time constant of adaptation reduced compared to wildtype littermates.

Conclusion

Plastin1 is important for correct formation and maintenance of inner hair cell stereocilia and defects due to its absence may account for the threshold shifts seen in plastin1 KO mice.


Organ of Corti Vibration Measured in the Intact Living Mouse Cochlea

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Background

The current knowledge of the cochlear mechanical process comes mainly from the direct measurement of the basilar membrane vibration. To understand how sounds excite mechanotransduction channels of sensory hair cells, it is critical to measure the vibration near the channels.

Methods

The sub-nanometer vibration of the organ of Corti was measured in sensitive mouse cochleae using a newly developed scanning heterodyne low-coherence interferometer (SHLI). The object beam of the SHLI was focused on the reticular lamina of the organ of Corti through the intact round window membrane, basilar membrane and cellular structures of the organ of Corti. The vibration magnitude and phase were measured as a function of frequency at different sound levels. The reticular lamina location was determined by measuring the tissue reflectivity as a function of the transverse location.

Results

At low sound levels, the vibration magnitude increased with frequency and reached the maximum at the best frequency. As the sound level increased, the response peak shifted toward low frequencies and the peak magnitude saturated. In sensitive cochleae, the organ of Corti vibration showed high sensitivity, sharp tuning and strong nonlinearity. Upon the death, the vibration magnitude decreased dramatically across different frequencies and showed broad tuning and the
linear growth. Compared to the basilar membrane vibration, the organ of Corti vibration showed higher sensitivity and stronger nonlinearity with no significant difference in the peak frequency and phase.

**Conclusion**
The current data demonstrate the active mechanical response of the organ of Corti in mice. These results provide essential information for studying the molecular mechanisms of cochlear amplification using genetically modified mice.

**Auditory Responses to Stimulation at Soft Tissue Sites Before and After Fixation of Mobile Components of the Middle Ear**

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**Background**
A new mode of auditory stimulation, namely, Soft Tissue Conduction (STC), which complements air conduction (AC) and bone conduction (BC) has been recently described. Since STC stimulation uses the same bone vibrator as in BC, it is possible that STC is simply a form of BC. The present study was conducted in order to investigate this possibility.

**Methods**
Six sand-rats initially underwent ablation of the left cochlea and opening of the right bulla. This was followed by measurement of auditory nerve and brainstem responses (ABR) thresholds to AC, BC and STC stimuli. AC stimuli were given through an insert earphone. BC through a bone vibrator to the skull and STC by applying the same bone vibrator to the sub-mental area. After baseline threshold measurements, the middle ear ossicles including the stapes footplate and the round window were totally fixated by immobilizing them with super glue (cyanoacrylate), thus eliminating the classical BC mechanisms. After the glue had dried, the AC, BC and STC ABR thresholds were determined again in the presence of the immobilization and compared to baseline measurements.

**Results**
Initial mean ABR thresholds before and after immobilization of the round window and ossicles in response to AC stimulation were 53.3±6.8 and 91.7 ± 11.3 dB pe SPL respectively. In response to BC stimulation the thresholds were 81.7±6.8 before and 80.8 ± 7.4 dB (instrument settings) after. With STC stimulation at the sub-mental area ABR thresholds were 99.2±2.0 before and 98.3 ± 4.1 dB (instrument settings) after the immobilization. Thus, mean air conduction responses were significantly elevated by 38.4 dB (two tailed paired t-test, p<0.00001) (conductive hearing loss), but mean BC and STC thresholds were virtually unchanged following the immobilization (in each, mean thresholds improved non-significantly by less than 0.9 dB; BC p= 0.61, STC p=0.36).

**Conclusion**
Even though the two windows were no longer mobile and bulk fluid flow is therefore no longer possible, and a pressure difference across the basilar membrane would not be induced, STC and BC thresholds were not altered. These and additional results from previous experiments provide evidence that STC may not be variant of BC, and that STC activation at low sound intensities may not be based on a passive traveling wave which requires two mobile windows.
Effect of Exposure to Intense Tones on Otoacoustic Emissions and Neural Thresholds
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\textbf{Background}
When the ear is stimulated with a pure tone the cochlea emits an “echo” at the frequency of the tone, so called stimulus-frequency otoacoustic emissions (SFOAEs). It has been hypothesized that SFOAEs at low probe levels originate within the peak of the traveling wave on the basilar membrane. This implies that SFOAE should carry place specific information about cochlear function and should be especially sensitive to small changes in the gain of the cochlear amplifier as compared to other type of emissions, specifically distortion product (DP) OAEs. Here we tested this prediction by recording SFOAEs and DPOAEs in chinchillas’ ears before and after exposure to an intense tone that resulted in clear shift compound action potential (CAP) thresholds.

\textbf{Methods}
All measures were obtained in anesthetized male chinchillas. The SFOAEs were measured for low level probes (30 dB SPL) varied from 0.3 to 12 kHz in fine steps with the suppression method. The suppressor was fixed near the probe frequency at moderate level. Additionally, SFOAE were measured with a second, high-frequency suppressor present (fs/fp ratio 1.2 – 2.5) to eliminate confounding effects of possible contributions from generators basal to the place of the probe frequency. DPOAEs were measured across a wide (f2=0.5-20 kHz frequency range (35/50 dB SPL, f2/f1=1.2). Animals were exposed to an intense tone (3 kHz, 100 dB SPL) to produce ~40 dB CAP thresholds shifts near 4 kHz and OAE tests were repeated.

\textbf{Results}
Even though in most cases SFOAE levels dropped following the exposure at frequencies where changes in CAP thresholds were evident, the SFOAE level shifts were always smaller and poorly correlated with amount of the shift observed for CAP. Measuring SFOAEs using the second high-frequency suppressor usually did not improve the match between SFOAE and CAP levels shifts. DPOAEs usually demonstrated a larger drop in level than SFOAEs that often better mapped the CAP threshold shift. However a variety of cases were noted ranging from DPOAE under- as well as over-estimating CAP threshold shifts.

\textbf{Conclusion}
SFOAE do not appear to be a more sensitive and accurate tool for evaluating changes in cochlear amplification as compared to DPOAEs measured with low stimulus levels.

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Transient Evoked Otoacoustic Emissions in Ears Exposed to an Intense Tone
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\textbf{Background}
It is commonly believed that otoacoustic emissions (OAEs) evoked with low level transient stimuli (TEOAEs) arise over a region of the basilar membrane mapped to the frequency content of the stimulus. This implies that TEOAE evoked with a narrow-band stimulus(tone-pip) should a) not contain energy outside of the band of the stimuli, b) be sensitive to changes in cochlear function only at frequencies within the stimulus band. We evaluate these predictions by measuring TEOAEs in chinchillas before and after exposure to an intense tone causing localized elevation in neural thresholds (CAP).

\textbf{Methods}
All measures were obtained in anesthetized male chinchillas. TEOAE were measured with the compression method using tone-pips (1 ms) centered at 1, 4 and 12 kHz. The stimulus level was fixed at 50 dB peSPL except few cases where additional recordings with lower levels where collected. Localized damage to the cochlea was induced with exposure to 3 kHz tone at high level to produce ~40 dB CAP threshold shifts at 4-6 kHz. Emissions were recorded before and after the exposure.

\textbf{Results}
For 1 kHz stimuli, TEOAEs showed small changes post-exposure, with the largest changes at the shortest delays. For 4 kHz stimuli, the spectra of TEOAEs pre-exposure usually had a frequency content similar to stimulus; but, an additional band centered around 0.5 kHz was usually present even for TEOAEs evoked with lower-level tone-pips. This low-
frequency band was particularly sensitive to acoustic trauma. The TEOAE centered around 4 kHz showed less consistent changes, typically a drop in amplitude with or without a change in fine structure. The 12 kHz emissions showed reduced amplitude only if CAP thresholds were elevated at the highest frequencies.

Conclusion
Given large threshold shifts near 4 kHz, the lack of consistent changes in the spectra of TEOAE evoked by 4 kHz tone pips near the center frequency of the stimulus suggests that the emissions originate partly in locations remote to 4 kHz place. But the 0.5 kHz band seems to represent an intermodulation distortion generated near the 4 kHz place since its amplitude was clearly reduced post-exposure. Changes to TEOAE fine structure as well as observed enhancements in amplitudes, particularly for 1 kHz stimuli, suggest that TEOAEs generally represent contributions from sources distributed over a wider region of the basilar membrane than hypothesized.

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Endocochlear Potential Powered Electronics
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Background
The endocochlear potential (EP) is a +70-100 mV electrochemical gradient generated between the intermediate and marginal cells of the stria vascularis and found within endolymph of the scala media. The EP provides approximately half of the electrochemical force driving auditory transduction currents and is critical to the cochlea's impressive sensitivity and dynamic range. Despite its role in powering active processes of the inner ear, the EP has never before been used as a power source for electronic devices.

Methods
Surgical exposure of a guinea pig cochlea was performed. The bulla was opened to provide access to the round window (RW) while maintaining the integrity of the tympanic ring and middle ear ossicles. Pulled glass micro-electrodes were advanced through the RW to access the fluid spaces of the inner ear. The negative electrode was inserted into the perilymph-filled scala tympani and the positive electrode through the RW and BM into the endolymph-filled scala media. Electrodes were connected to a newly designed, anatomically-sized, ultra-low quiescent-power energy harvester integrated with a wireless transmitter (endo-electronics chip). The impact of experimental procedures on the physiological function of the ear was assessed via tone-pip audiogram and measurement of the compound-action potential (CAP).

Results
The endo-electronics chip was successfully operated for periods of up to 5 hours while utilizing a guinea pig EP as the power source. During this time the EP remained stable, with no decline indicative of long-term damage from over-taxing the energy supply. CAP testing performed before and immediately after electrode insertion demonstrated that electrodes could be placed without causing hearing impairment. Comparison of CAP results before and after prolonged current draw similarly confirmed that power extraction is possible without sacrificing hearing performance.

Conclusion
We have demonstrated that it is possible to utilize the mammalian EP as the source of energy for electronic devices, in this case a radio-transmitter capable of broadcasting sensory information to external devices. This has been accomplished without significantly compromising the ear's ability to transduce sound. With improvements in electrode design, this biologic battery may provide novel opportunities to sustainably power sensors or drug-delivery actuators in the vicinity of the ear, eye and brain of mammals; enabling transformative opportunities for new diagnostics and therapies in hearing research and related fields.
Endolymphatic microphonics to low-frequency tones and noise, and their suppression by higher-frequency sounds: A study relevant to wind turbine noise.

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Background
Large cochlear microphonics (CM) recorded from scala media have been reported with infrasonic fluid-injected stimuli (Salt and DeMott, JASA 106:847; 1999). In the current study we investigate endolymphatic CM responses with a variety of low-frequency acoustically-delivered stimuli.

Methods
DC-coupled CM recordings were made from endolymph of the first and third turns of guinea pig cochleae in response to tonal and low-pass filtered frozen noise stimuli.

Results
In response to tonal stimuli, the maximum CM amplitude – the peak of the growth curve as stimulus level was varied – increased markedly as frequency was lowered. Specifically, the maximum CM amplitude averaged 4.0 mV at 500 Hz (n=9), 10.8 mV at 50 Hz (n=11), and 17.2 mV at 5 Hz (n=11). The large CM to 5 Hz stimulation was suppressed by the simultaneous presentation of a 500 Hz stimulus. A similar suppression of low-frequency responses was observed when microphonic was evoked with low-pass filtered noise and the filter cutoff frequency was increased. CM, measured as a spectral average across the 12-125 Hz range with the low pass noise cutoff frequency at 125 Hz, averaged -70.1 dB re 1V (n=5). As the filter cutoff frequency was increased in half-octave steps, CM responses across the 12-125 Hz range decreased progressively, reaching -78.7 dB re 1V (n=5) with the cutoff filter at 1.4 kHz.

Maximum endolymphatic CM amplitude from the first cochlear turn in response to tones and low-pass filtered noise was lower in amplitude than in the third turn. Additionally, noise-evoked measures showed smaller changes as the filter cutoff frequency was varied.

Simultaneously recorded output from a microphone in the ear canal was virtually unchanged as the noise-stimulus filter was varied, demonstrating that the changes to endolymphatic CM did not result from variations in signal power.

Conclusion
These data show that in the absence of higher-frequency sounds, endolymphatic responses to infrasonic stimuli can be substantially larger than to stimuli in the audible range. These large responses are relevant to situations where people are exposed to very low-frequency sounds when ambient noise levels are low. This occurs within people’s homes located near wind turbines. Although the house structure attenuates the higher frequency sounds, it does not attenuate the infrasound generated by the wind turbine to the same degree. If the ear responds strongly under conditions such as this, it may account for sleep disturbances and other symptoms that some living near wind turbines report.
Logic of sound level encoding in the auditory nerve

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1Inserm

Background
The mammalian cochlea encodes sound level over a large dynamic range, through the progressive recruitment of three auditory nerve fibers pools, the high-, medium- and low- spontaneous rate (SR) fibers, with different threshold and dynamic range.

Methods
Using a combination of pharmacological, anatomical and computational approaches, we investigate the weight of each auditory fibers pool on the sound-evoked compound action potential (CAP), as a proxy of sound level dynamic range.

Results
Our results demonstrate that CAP thresholds and amplitude are independent of the low-SR fibers but rely on the high- and medium-SR fibers. This argues against a continuum activation of all the auditory nerve fibers to encode the full sound-level dynamic range. In addition, we suggest that intrinsic properties of high-SR fibers suffice to explain CAP dynamic range.

Conclusion
Our results show that substantial auditory nerve fibers loss can co-exist with normal hearing and call for relevant clinical tests to detect hidden auditory nerve fibers degeneration.

Spiral Ganglion Myelination across Mammalian Species

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Background
Only human lacks a myelin sheet around the soma. The meaning for this difference remains unclear and may be related to facilitate synchronous activity or preprocess the auditory signals.

Methods
We performed a comparative and quantitative analyses of the spiral ganglion in various species including humans, monkey, pig, sheep, guinea pig and mice. Further we analyzed the structure of monopolar spiral ganglion neurons (SGNs) in human in cases of noise induced hearing loss and Wolfram disease in human and compare this with SGN degeneration in animal models.

Results
Results show that human type I neurons have cell bodies that are umyelinated to 90-97%. The type II cells making up approximately 3-5% are completely unmyelinated in mammalian species including peripheral and central axons. The unmyelinated nature in human is true for at all frequency levels. Findings highlight the marked morphological differences in the structure of the spiral ganglion between human and other mammalian species.

Conclusion
The presence of intimate contacts between spiral ganglion neurons in man around their perikarya supports the hypothesis of transcellular communication. The lack of a rigid myelin layer may also elucidate the healing mechanism when peripheral axons are lost retrograde to hair cell death and explain the robust survival of monopolar SGNs.
HUMAN SPIRAL GANGLION IS UNMYELINATED – WHY IS SO?
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Background
Thorough morphologic analysis of the human spiral ganglion obtained after direct fixation of surgical specimens in normal hearing individuals show that they are unmyelinated contrary to animals studied so far. The reason for this difference remains obscure but may be essential both for understanding the normal physiology and pathology especially after sensory deafferentiation.

Methods
In here we analyzed systematically the structure of the human spiral ganglion at various frequency levels in optimally fixed specimens from normal hearing individuals undergoing petroclival meningioma surgery. Verification of the normal hearing is demonstrated by audiometry. The inner ear as well as the auditory pathways were unaffected by disease.

Results
Results show that human type I neurons have cell bodies that are unmyelinated to 90-97%. Central and peripheral axons are myelinated in 100%. The type II cells making up approximately 5% are also completely unmyelinated including peripheral and central axons. The unmyelinated nature is verified at all frequency levels. In addition the unmyelinated type I cells often form cellular clusters with intimate contacts between somata. At these places separating satellite glial cells is often incomplete forming direct contacts between individual cell body surface membranes. This may allow for electric cross excitation and facilitate trans-cellular communication between cells at the level of the ganglion.

Conclusion
Findings highlight the marked morphological differences in the structure of the spiral ganglion in different species. The exceptional conditions of the human spiral ganglion raises questions about the possibility of generating specific complex and dynamic auditory responses essential for human speech communication. The conditions may also play an important role for the unique preservation properties of the human spiral ganglion cells following sensory hair cell loss.
Cooperative Release of Synaptic Vesicles at a Hair Cell Ribbon Synapse

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Background
According to the classic quantal theory of synaptic transmission in the NMJ, each release site (or active zone) has a low release probability and can release only one synaptic vesicle in response to one presynaptic action potential. In addition, the releases of synaptic vesicles are independent from each other. However, synchronized release of a few synaptic vesicles from the same release site, i.e., multivesicular release, has been demonstrated at hippocampal synapses, retinal bipolar cell ribbon synapses and hair cell ribbon synapses. At bipolar cell ribbon synapses the release of synaptic vesicles has been shown to be coordinated and binomial, but not cooperative. At hair cell synapses, however, the spontaneous multivesicular release events have been suggested to be cooperative. In the present study, we investigated whether synaptic vesicle releases from frog auditory hair cells are cooperative during physiologically relevant stimulations.

Methods
Paired patch-clamp recordings were made on hair cells and their connected afferent fibers in bullfrog amphibian papilla. Hair cells were stimulated with sinusoidal voltage-clamp commands (400 Hz, with 5, 10 and 20 mV peak-to-peak amplitudes, centered at -55 mV), which mimic hair cell voltage responses in vivo to pure sound tones. Excitatory Postsynaptic Currents (EPSCs) were recorded from afferent fibers, and individual synaptic events were detected with a sliding template technique. Based on our previous study on single vesicle (or quanta) events, the quantal contents were calculated for the evoked EPSC events.

Results
By varying the strength of stimulation, we found that although the failure rate of EPSCs can be as high as ~40%, the majority of the EPSC events still contains 2~3 vesicles. A simple binomial model cannot account for this finding. However, the data can be fit well with a simple cooperative release model where the release of one synaptic vesicle increases the probability of nearby vesicles being released, thus producing a nearly synchronous multivesicular event.

Conclusion
The release of synaptic vesicles from auditory hair cells under physiological stimulations is cooperative and cannot be described by a simple binomial distribution.
Hair Cell and Neural Potentials Recorded at the Round Window in Gerbils
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Background
Electrocochleography measured at the round window contains both hair cell and neural responses. Responses from the hair cells are the cochlear microphonic (CM) and summating potential (SP). Responses from the auditory nerve are the compound action potential (CAP) and auditory nerve neurophonic (ANN). The CAP and SP are relatively isolated and can be readily identified in the waveform. The CM and ANN are both present in the ongoing part of the response; the CM follows the fine structure of all frequencies and the ANN is the evoked response correlate of phase locking in the auditory nerve. Because these signals are mixed, it is a challenge to separate them.

Methods
ECoG potentials were measured from the round window of Mongolian gerbils to tones of different frequencies and intensities. Forward masking was also used in an attempt to attenuate the neural response. Gerbils were either normal hearing or had a pattern of high frequency noise induced hearing loss (NIHL) similar to many cochlear implant recipients.

Results
In normal hearing gerbils, distortions in the ECoG waveform, likely attributable to phase locking associated with the ANN, were present to low frequencies at low and moderate intensities. At high intensities, distortions were present at all frequencies, suggesting saturation of stereociliary movement. Forward masking with a masker 30 dB louder than the probe reduced the signal and distortions across the frequency range including high frequencies, suggesting that a preference for closed channels in stereocilia persists for a time after the stimulus. In gerbils with NIHL, the ongoing part of the ECoG saturated at relatively low intensities indicating a loss of spread of excitation.

Conclusion
These results show that distortions in the ECoG attributable to phase locking in nerve fibers are restricted to low frequencies at low and moderate intensities, and that forward masking is present in hair cells at high intensities.
Firing patterns of afferent calyces differ with epithelial zone in the immature rat saccule

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Background

The mammalian vestibular epithelium has unique calyceal synapses between type I hair cells and the postsynaptic afferent terminal. Two classes of vestibular afferents form calyceal endings: calyx-only fibers innervate the central striolar zone of the epithelium, and dimorphic fibers form both calyceal and bouton endings on hair cells and innervate either the striola or the extrastriola. Striolar afferents have irregular spike timing whereas extrastriolar afferents have regular spike timing. From work on isolated vestibular ganglion cell bodies, irregular spiking is thought to correlate with the transient responses to current steps, and regular spiking with sustained responses. Here we investigate these proposed relationships with whole-cell recordings from striolar and extrastriolar calyces in the epithelium, where spiking initiates.

Methods

Whole-cell current clamp recordings were made from calyces in a semi-intact preparation of the immature (postnatal days 2-9) rat saccular epithelium and attached nerve. Nomarski optics and fluorescent dye fills were used to identify the morphology of the afferents.

Results

Striolar calyces responded to depolarizing current steps with a single spike at the stimulus onset, consistent with recordings from vestibular ganglion. At 25-29°C, 19 of 24 calyces were silent, and the remaining five had low and irregular spontaneous activity: 6±2 spikes/s, CV=1.14±0.25, Fano factor=0.24±0.06, n=5. In contrast, most extrastriolar calyces gave sustained responses to depolarizing current steps (12/16). They were also more likely to fire spontaneously (9/16) and the firing was higher-rate and more regular: 15±2 spikes/s, n=9 p<0.014; CV=0.22±0.08, p<9e-4; Fano factor=0.013±0.008, p<1.8e-4.

Firing rates are expected to be higher at body temperature. Indeed, in 7 complex calyces recorded at 35-37°C, four fired spontaneously, with higher firing rates than recorded at room temperature (thermal Q10 of 3).

The spontaneously active striolar calyces were from P7 or P8 animals, while many of the silent calyces were from P2-P4 animals, raising the possibility that spike activity increases with maturation.

Conclusion

Striolar calyces had transient responses and irregular spontaneous activity. Extrastriolar calyces had sustained responses and regular spontaneous activity. These results suggest that in vivo differences in spike regularity between the striolar and extrastriolar zones are preserved in our preparation, and are consistent with evidence correlating specific ion channels with firing patterns and spike timing.
Discharge Regularity and Information Transmission in the Mammalian Crista
Jay M Goldberg

University of Chicago

Background
Two themes are common in vestibular organs. 1) Regular and irregular afferents exist side-by-side. 2) The two kinds of afferents differ in their response dynamics. Can a rationale be provided for these two themes? In particular, can we explain why there is a close association between response dynamics and discharge regularity despite the fact that there is both experimental and theoretical evidence that these two discharge properties are not causally related? One possibility is that the association is needed for the efficient encoding of sensory information.

Methods
Efficient encoding can be measured by the mutual information between stimulus and response. For this reason, we calculated upper-bound estimates of mutual information (MIUB) at several frequencies (f). MIUB is proportional to the square of the ratio between $G(f)$, the conventional rotational gain (in spikes per s/deg per s) and $CV$, the coefficient of variation of the background discharge. Gains and CVs were obtained from Ramachandran and Lisberger (2008) over a frequency range from 0.5-10 Hz.

Results
Reflecting a flat gain and a low CV, the MIUB of regular units is uniformly high throughout the frequency spectrum. High-gain irregular (dimorphic) units are characterized by a high CV and a frequency-dependent gain increase; the result is an MIUB with an upward slope that crosses the regular curve near 1 Hz. Low-gain irregular (calyx) units are especially phasic; given their low gain, high CV and phasic response dynamics; their MIUB curve rises steeply, but only crosses the regular curve at frequencies approaching 20 Hz.

Conclusion
Regular and high-gain irregular units provide more efficient coding at low and high frequencies, respectively. Low-gain irregular units may be specialized to handle rapid head saccades with peak head velocities approaching 500 deg/s. The superior performance of irregular units at high frequencies depends on a 10-fold frequency-dependent gain increase, which be achieved only by the high sensitivity to depolarizing inputs of the hair cells synapsing on such units. This same characteristic is needed to produce an irregular discharge. A need for high synaptic sensitivity may explain the association between an irregular discharge and phasic response dynamics.
The Anatomy of the Lamprey Ear: Morphological Evidence for Horizontal Canal Equivalent in the Labyrinth of Lamprey, Petromyzon marinus

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Background

Jawed vertebrates’ (Gnathostome’) inner ears have three semicircular canals arranged orthogonally in the three Cartesian planes: one horizontal (lateral canal) and two vertical canals each with its own ampulla containing a canal crista uniformly polarized toward (horizontal crista) or away (vertical cristae) from the utricle. Canals with cristae function as a detector for angular acceleration movements in their respective planes. In jawless craniates, cyclostomes (hagfish and lamprey), the horizontal canal is missing. The hagfish inner ear has one doughnut shaped torus (equivalent to the two vertical canals) vertical canal, and the lamprey has reportedly only two vertical canals. These observations on the anatomy of the cyclostomes’ inner ear have been unchallenged for over a century, and the question of how these jawless vertebrates perceive angular acceleration in the horizontal (yaw) plane remains open.

Methods

In the present study we sought an answer to this question by reevaluating the anatomy of the inner year in the sea lamprey, Petromyzon marinus, using stereoscopic dissection and scanning electron microscopy.

Results

In contrast to all gnathostomes, the lamprey has two horizontal canals in each labyrinth. In contrast to jawed vertebrates, the horizontal canals in lampreys are located on the medial surface in the labyrinth rather than on the later surface. These radical differences suggest that the horizontal canal’s evolved twice and independently as a lateral canal associated with only one crista in gnathostomes and as two symmetric medial demi-canals, each associated with the horizontal leg of the tripartite canal ampulla.

Conclusion

Given that the sensory epithelia of the canal cristae in hagfish, lampreys and gnathostomes differ in structure and distribution and that lampreys and hagfish apparently are a monophyletic taxon, it is most parsimonious to assume that each canal system with associated cristae evolved independently from proto-chordate precursors ears that likely had no canals or cristae but served only as gravistatic (statocyst) sensors as in most aquatic non-vertebrates.

Differences in the Diameter of Facial Nerve and Facial Canal in the Pathophysiology of Bell’s Palsy – A Three-Dimensional Temporal Bone Study.

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Background

Bell’s palsy is hypothesized to result from virally mediated neural edema. Ischemia occurs as the nerve swells in its bony canal, blocking neural blood supply. Since viral infection is relatively common and Bell’s palsy relatively uncommon, it is reasonable to hypothesize that there are anatomic differences in the diameter of the facial nerve canal that predispose some patients to develop paralysis. Measurements of the facial nerve and canal as it follows its tortuous course through the temporal bone are difficult without a 3D view. In this study, 3D reconstruction was used to compare anatomic differences in temporal bones of patients with and without a history of Bell’s palsy.

Methods

Of 78 patients that presented with facial paralysis from the human temporal bone (HTB) collections of the University of Minnesota and MEEI, 6 patients met diagnostic criteria for Bell’s Palsy (BP): 5 unilateral and 1 sequential bilateral case. Three-dimensional models were generated from HTB histopathologic slides with reconstruction software (Amira®), and the diameters of the facial canal (FC) and facial nerve (FN) were measured at the midpoint of the labyrinthine, tympanic, and mastoid segments. Measurements were compared between HTB with BP to the contralateral, unaffected side of each case (CLS) and to 10 age-matched normal controls using non-parametric tests.

Results

The mean diameter of the FC and FN was significantly smaller in the tympanic and mastoid segments (p=0.01) in the BP group than in the control group. There was no difference between groups in regards to the diameter of the labyrinthine segments (fig.1). The FC to FN diameter ratio (FC/FN) was smaller in each of the three segments in the BP group.
compared to controls, but the difference was statistically significant only in the mastoid segment (fig.2). There were no significant differences between the diameters of the FC or FN between the BP and contralateral sides. When comparing the BP and control groups, the narrowest part of the FC was the labyrinthine segment in the control group and the tympanic segment in the BP.

**Conclusion**
This study suggests an anatomical difference in the diameter of the FC in the tympanic and mastoid segments but not in the labyrinthine segment in patients with Bell’s palsy. These findings suggest that there may be a site of anatomical predisposition in the FC that may have treatment implications.
Developing a mouse model for cochlear implantation
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Background
Animal models are the only means of assessing the effects of cochlear implantation (CI) at a cellular and molecular level. Mice are an excellent model for auditory research and this is in part due to the range of naturally occurring and genetically-modified strains which mimic human deafness. To date, there are very few studies of CI in mice. Our aim was to create a mouse model of CI and subsequently assess the effects of CI on cochlear ultrastructure and function.

Methods
Surgical techniques were developed and refined using cadaveric dissection prior to live surgery. C57Bl6 mice (early-onset hearing loss model) were the initial CI group. Access to the round window was gained and implantation using a specialised electrode array (n=6) and dummy implant (n=12) performed. The contralateral cochlea acted as a control. Auditory brainstem response (ABR) audiometry prior to and at time-points following CI was undertaken. Cochleae were fixed and decalcified before being prepared for histological examination.

Results
All mice recovered well following surgery. Postoperative ABR testing revealed a mean elevation in thresholds of 10dB when compared to preoperative levels. This was a permanent threshold shift in the majority of cases. Correct placement of the electrode array was confirmed using cone beam CT and light microscopy. Initial analysis revealed encapsulation of the implant in tissue with features suggesting the presence of fibrosis. Immunolabelling using a leucocyte marker, CD45, revealed greater numbers of positively labelled cells within the basal turn of implanted cochleae compared to controls. This was particularly seen within the region of the spiral ligament and basilar membrane, suggesting enhanced recruitment of these cells.

Conclusion
Our results show that mouse CI via the round window is achievable and holds many translational benefits. Patient studies have suggested that fibrous encapsulation reduces implant efficacy, prompting the necessity for replacement in some. Further investigation into the presence of inflammation and fibrosis using this mouse model will therefore help identify measures to potentially reduce these effects. The model will allow for future research using eluting electrode arrays as a means for drug delivery to cochleae, ultimately to enhance patient outcomes.

Preservation of Residual Low Frequency Hearing in a Guinea-Pig Model of Non-Electrically Active Cochlear Implantation.
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Background
Electro-acoustic stimulation (EAS) with Cochlear™ Hybrid implants is a same-ear bimodal concept that has extended the traditional implantation candidacy criteria to patients that have retained some functional hearing in the low frequencies. Hearing loss in such candidates is targeted with electrical stimulation of the basal cochlear turns, encoding the high frequencies, in concert with acoustic stimulation of the low frequency apical turns. However, the perceptual benefits
gained through EAS are hampered by clinical evidence that some patients eventually lose a considerable amount of residual hearing in the implanted ear. The aetiology of this loss of low-frequency sounds is currently unknown; an inflammatory-fibrotic 'foreign body' response triggered post-implantation (PI) is one mechanism thought to be involved. To identify the molecular mechanism(s) underlying this loss of low-frequency hearing, we have established an animal model of residual hearing in the guinea-pig and developed a soft surgical approach for cochlear implantation via the round window.

**Methods**
Guinea-pigs were treated with gentamicin (125mg/kg) for 10 consecutive days, a procedure in our laboratory that has consistently shown to produce hair cell loss restricted to the basal turns. Auditory brainstem responses (ABRs) to click and tone pip stimuli (4-40 kHz) were recorded to confirm residual low frequency hearing. Guinea-pigs were implanted with a 14-electrode array. Additional ABR recordings were taken pre-implant and monitored until sacrifice: 2 days, 1 week and 1 month PI. Auditory bullae were fixed and tissue prepared for histological processing.

**Results**
Little elevation in ABR thresholds was evident in implanted ears 2 days PI compared to contralateral control ears. At 1 week PI threshold shifts of 10-20dB were identified in implanted ears compared to control, that exhibited minimal further deterioration in hearing up to one month PI. At 2 days PI, a soft tissue covering the electrode array was observed, which at 1 week PI had expanded to encapsulate the electrode array. One month PI, tissue consistent with the appearance of neo-osteogenesis was observed along the implant pathway externally from the round window. Immunofluorescence analysis with an anti-CD45 antibody showed enhanced immunoreactivity in the scala tympani of implanted ears compared to controls; CD45 positive cells consistent with the morphology of macrophages were identified from 2 days PI.

**Conclusion**
Threshold shifts following cochlear implantation manifest within 1 week PI in a guinea-pig model of residual hearing, which is consistent with the timing of fibrosis observed in the scala tympani over the electrode array.
Novel In Vitro Model for Cochlear Electrode Induced Mechanical Trauma Studies
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Background
Mechanical trauma from cochlear implant electrode insertion can lead to inflammation and oxidative stress reactions within the cochlea that result in the apoptosis of the auditory hair cells (HC) and initiation of intracochlear fibrosis. To gain insight into the molecular mechanisms that can cause this trauma initiated loss of auditory HC and cochlear fibrosis and the otoprotective effects of dexamethasone (DXM) observed in our previous in vivo studies, an EIT in vitro model was created and characterized.

Methods
Cochleae from 3 days rat pups were divided into: 1) Control untreated; 2) EIT; 3) EIT+ DXM (50 µM). In groups 2 and 3, a 0.3 mm diameter monofilament fishing line was introduced through a small cochleostomy performed next to the round window membrane, in order to achieve a high angle of insertion, i.e. between 110-150º. Organ of Corti explants were dissected and then either untreated or treated with DXM. HC counts, gene expression for pro-inflammatory cytokines (i.e. TNFα and IL-1β), pro-inflammatory inducible enzymes (i.e. iNOS and COX-2) and growth factors (i.e. TGFβ1, TGFβ3 and CTGF), oxidative stress (i.e. CellROX), and analyses of apoptosis pathways (i.e. Caspase-3, AIF and Endo G) were carried out on all explants.

Results
Expression levels of both TNFα and IL-1β inflammatory process related genes increased significantly in response to EIT and these increases were prevented by DXM treatment. EIT in this in vitro model also caused a significant increase in oxidative stress and loss of HCs that was prevented by treatment with DXM. EIT-induced HC losses were caused by conversion of procaspase-3 to its active form. The strongest effects of these EIT-initiated changes were observed mainly in the basal and middle turns of the EIT explants.

Conclusion
This new EIT-model of insertion trauma provides a useful tool for understanding the mechanisms involved in both inflammation-induced HC losses and the initiation of fibrosis that can occur following cochlear implantation.
Cochlear electrode implantation trauma causes over-expression of TGF-β1 and activation of the Wnt/β-catenin pathway in an in vitro model

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Background

The insertion of a CI electrode array can be associated with an inflammatory response that is known to induce loss of auditory hair cells and growth of fibrotic tissue around the electrode array. This fibrotic process can have a negative impact on the function of a patient’s implant. Wnt/β-catenin/Tcf signaling induces the expression of Axin2, which in turn participates in a negative feedback loop to limit the intensity of a Wnt-initiated signal. From previous work we know that trauma to the cochlea induces the expression of TGF-β1 and we hypothesize that this growth factor can co-operate with the Wnt pathway to elicit fibrosis.

Methods

Cochleae from 3 days rat pups were divided into: 1) Control untreated; 2) TGF-β1; 3) TGF-β3; 4) EIT; 5) EIT+ TGF-β1 (10 ng/ml); and 6) EIT+ TGF-β3 (10 ng/ml). qRT-PCR for Axin2; Wnt4, Wnt5b and CTGF was performed in organ of Corti (oC) explants and lateral wall tissues (LW) after 96 hours of incubation.

Results

Upon EIT, CTGF was up-regulated in oC and LW tissues, compared to control explants. Addition of TGF-β1 and TGF-β3 to the LW tissues resulted in the upregulation of CTGF. In the EIT-TGF-β3 oC explants the CTGF levels were as low as the control group. Axin2 mRNA levels were remarkably increased in oC explants either treated with TGF-β1 or TGF-β3. Wnt4 was downregulated in the LW tissues by the addition of TGF-β1 and TGF-β3, but no differences were observed between control and EIT groups. While Wnt4 levels were not altered in the oC explants, a decrease in Wnt5b mRNA levels was observed upon EIT, and in groups 2 and 3 compared to 1. Interestingly Wnt5b was upregulated in the EIT group treated with TGF-β3.

Conclusion

Induction of EIT in the cochlea triggers a fibrogenic response as part of the wound healing process, in which TGFβs are key role players. oC cells and LW cells play different roles. While there is an increase on the Wnt/β-catenin activation pathway upon EIT in oC, exacerbated by TGF-β1 treatments, a downregulation of the Wnt5b was observed. This is consistent with previous reports that indicate antagonism between the canonical and noncanonical Wnt pathways. Our results suggest that the Wnt pathways do not participate in the fibrotic response initiated by the lateral wall tissues.
Age related changes in the inner ear of the mouse model of metabolic syndrome
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Background
The metabolic syndrome is characterized by obesity concomitant with other metabolic abnormalities such as hypertriglyceridemia, hyperlipidemia, elevated blood pressure and diabetes. It is known that the prevalence of hearing loss is high in the patients with diabetes. However, the causes of hearing loss have not been clear. In the present study, we evaluated the hearing function in animal model of diabetes with obesity.

Methods
TSOD male mice used in study were derived from the Tsumura Research Institute. Age- and sex-matched non-diabetic TSNO mice served as controls. Blood samples were collected from TSOD and TSNO mice at 3, 5, 8 months of age. Blood glucose concentrations of TSOD mice were significantly higher than those of controls. In addition, Body weights of TSOD mice were greater than those of controls.

Results
We evaluated the ABR thresholds of animals in 3, 5, 8, 12 month of age. There was no different in ABR thresholds between TSOD mice and controls in 3 or 5 month of age. However, the elevation of ABR thresholds were observed in 8 month aged TSOD mice. The thresholds of 8 months aged TSOD mice were significantly larger than those in controls. The histological examination showed the thickened vessel wall of cochlear modiolus, and the arterial stenosis in the stria vascularis. In addition, the gene expression profiles were analyzed using the Affymetrix GeneChip®. The microarray analysis indicated the expression of growth factors such as IGF-1 in the model mouse related with control mouse.

Conclusion
The understanding of changes in the inner ear of the model animals might be of importance in future prophylactic approaches to prevent hearing loss in the patient with metabolic syndrome.

A novel animal model for studying blast injuries of the inner ear using laser-induced shock waves.
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Background
Recently, the number of blast injuries of the inner ear has increased in frequency in the general population. Inner ear injury may result in the temporary or permanent sensorineural hearing loss, but their mechanisms underlying this phenomenon are not fully understood. The majority of animal models for these injuries use explosive or complex...
experimental settings, limiting the laboratory study of blast injury. Recent studies have revealed that brain and pulmonary injuries caused by laser-induced shock waves (LISW) would be useful models to study blast injury.

**Methods**
The aim of this study was to establish a small-animal model for inner ear injuries associated with blast injury, using LISW with high controllability, and easy experimental settings. A LISW was generated by the irradiation of an elastic laser target with 694nm nanosecond laser pulses of a ruby laser. Anesthesia was initially induced in Sprague-Dawley rats (5w). After transmastoidal cochlear laser-irradiation at 6.6 J/cm², 7.3 J/cm², 8.3 J/cm² in rats, we investigated the auditory brainstem response (ABR) and distortion product otoacoustic emission (DPOAE) after the laser irradiation, and outer hair cells (OHCs) loss was examined using surface preparation technique. Immunohistochemical analyses for 8-hydroxy-2-deoxyguanosine (8-OHdG) were performed to examine the amount of oxidative DNA damage that occurred in cochlea.

**Results**
ABR and DPOAE measurements revealed that this animal model has the auditory dysfunction associated with inner ear damages. With regard to the peak pressure of laser-induced shock wave, there was a better improvement for the 6.6 J/cm² group as compared to the 8.3 J/cm² group. Furthermore a significant lower survival rate of OHCs was observed of the 8.3 J/cm² group as compared to that of 6.6 J/cm² group and strong immunoreactivities against 8-OHdG were observed in the inner ear tissues of the laser-irradiation rats. These data suggested that 6.6J/cm² group seems to be a temporary hearing loss (TTS) models and 8.3J/cm² group to be a permanent hearing loss (PTS) models.

**Conclusion**
Animals exposed to transmastoidal cochlear laser irradiation showed functional and morphological changes of the inner ear similar to other studies in blast injury. Although the underlying mechanism of blast injury seems to be complicated, this novel rat model of blast injury using LISW is suitable for detailed studies of blast inner ear injury in the laboratory.
Hearing impairments and cochlear changes in type1 diabetes model mice and diet-induced obesity mice.

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Background
Recent several population-based studies suggested the association between diabetes and hearing impairment. However, the impact of diabetes on hearing impairment has not been as well recognized in comparison with the known complications affecting the renal, visual, and peripheral nervous systems. In addition, the association between obesity and hearing loss also has been reported.

Methods
1. Type1 diabetes model
Streptozotocin induced male C57BL/6J diabetic mice were used. Hearing function was evaluated 1, 3 and 5 months after induction of diabetes (n = 10 per time point) using auditory evoked brainstem responses (ABR). Mice (n = 8) were exposed to loud noise (105 dB) 5 months after induction of diabetes. ABR were measured before and after noise exposure. Cochlear blood flows were measured by laser-Doppler flowmeter. Spiral ganglion cells (SGCs) were counted. Vessel endothelial cells were observed by CD31 immunostaining.

2. Diet-induced obesity model
C57B/6J and CBA/N male mice were used. Obesity was induced with a high fat diet (HFD32, CLEA Japan). Hearing function was evaluated 1, 3 and 5 months (B6, n = 10 per time point) and 1 year (B6, n=10 and CBA, n=9) after induction of high fat diet.

Results
1. Type1 diabetes model
Chronologic changes in the ABR threshold shift were not significantly different between the diabetic group and controls. However, vessel walls in the modiolus of the cochleae were significantly thicker in the diabetic group than the control group. Additionally, recovery from noise-induced injury was significantly impaired in diabetic mice. Reduced cochlea blood flows and SGC loss were observed in diabetic mice cochleae after noise exposure.

2. Diet-induced obesity model
At 5 month after induction of obesity, compared to the baseline, ABR threshold shifts were significantly increased in Control at 32kHz compared with Obesity model in B6 mice. Moreover, 1 year after induction, ABR thresholds were significantly increased in Control at all frequencies. However in CBA mice, ABR thresholds were significantly increased in Obesity model compared with Control at all frequencies at 1 year after induction.

Conclusion
Our data suggests that diabetic cochleae are more susceptible than controls to loud noise exposure, and decreased cochlear blood flow due to sclerosis of the vessels and consequent loss of SGCs is a possible mechanism of hearing impairment in diabetic patients.

Attempts to Rescue Outer Hair Cell Loss Induced by Dysfunctional Prestin in 499 Prestin Knockin Mice

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Background
Prestin is the motor protein responsible for outer hair cell (OHC) somatic electromotility. Removal of prestin results in a change in cochlear mechanics, as well as a loss of amplification. In order to study this process, a knockin (KI) mouse model was created to preserve physical/anatomical OHC properties, which are altered in prestin knockout (KO) mice. The resulting 499 prestin KI (Dallos et al., 2008) was created by replacing 2 of prestin’s 744 amino acids: V499G/Y501H. These mice have vastly reduced electromotility and sensitivity, but maintain normal OHC length and stiffness. Although 499 OHCs appear WT-like, hair-cell death is accelerated relative to that documented in mice lacking prestin protein.

Methods
Because the loss of OHCs complicates the interpretation of experimental results, we implemented three interventions. First, 499 KI’s were backcrossed to BAK KO mice (C57BL6), which lack the mitochondrial pro-apoptotic gene Bak, in the
hope of reducing apoptotic cell death. Second, because oxidative stress is a major cause of OHC death, another group of 499 KI mice was fed the antioxidant diet, Protandim (Liu et al., 2009). This approach in chemoprevention is thought to decrease the effects of oxidative stress by increasing the expression levels of superoxide dismutase, as well as catalase activities. Finally, 499 KI mice were backcrossed onto the FVB murine strain in order to avoid the relatively early onset age related hearing loss associated with the original 129/B6 499 KIs. Hair-cell loss was documented by constructing cytocochleograms at 42 days of age.

Results
A significant reduction in OHC death (Student’s t-test) was observed in FVB/499KI mice, and in mice receiving the Protandim diet, intimating an increased preservation of OHCs. 499 KI mice that were also homozygous for the Bak gene failed to show any improvement in OHC death.

Conclusion
Since the benefit obtained for FVB/499 KI mice was more dramatic than that achieved with Protandim, these results provide guidance for future interventions designed to ameliorate progressive OHC loss. (Work supported by NIDCD Grants DC00089 and DC010633, and the Knowles Hearing Center).

In-depth Proteomic Analysis of Mouse Cochlear Sensory Epithelium by Mass Spectrometry
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Background
Identification (ID) of the complete proteome in sensory organs such as the cochlea is challenging due to small amounts of tissue and because membrane proteins, such as ion channels, are difficult to isolate and identify. Several proteomic techniques have been developed, including 2D-DIGE, antibody microarray and mass spectrometry. However, mass spectrometry in conjunction with separation methods can provide a more comprehensive proteome because of its ability to enrich protein samples, detect low molecular weight and hydrophobic proteins, and identify low abundant proteins by reducing its dynamic range.

Methods
We used several separation techniques to maximize protein ID, applied Filter Assisted Sample Preparation (FASP) for detergent removal, and varied protein digestions prior to using nano liquid chromatography tandem mass spectrometry (MS), to obtain an in-depth proteome analysis of 30-day-old mouse cochlear sensory epithelium. In the first approach, we fractionated extracted proteins using gel-eluted liquid fraction entrapment electrophoresis (GELFrEE). In the second, proteins were digested and separated by strong cation exchange (SCX) or weak anion exchange (WAX) chromatography. We also compared single tryptic digestions with double tryptic or tryptic and LysC digestions.

Results
Double digestion with SCX fractionation identified 3390 proteins with a 1% false discovery rate (MaxQuant), of which 497 were in the plasmalemma and 19 of these were ion channel α-subunits. GELFrEE yielded 2173 proteins, of which 511 were absent when using SCX. Gene ontology annotation showed that 59% and 51% are binding proteins and 32% and 40% are catalytic proteins when comparing SCX and GELFrEE, respectively. GELFrEE identified 10 ion channel α-subunits with one unique channel not observed by SCX. The distribution of biological processes, cellular components, and protein classes were comparable between different separation methods. Combining the results from the different techniques resulted in 4187 total protein IDs, including 21 ion channel α-subunits. Additional protein data filtering using Scaffold produced similar ion channel IDs at 95% confidence, but with 19% higher numbers of identified channels.

Conclusion
The results show that the application of multiple approaches is needed to provide an exhaustive proteome analysis of the cochlear sensory epithelium that includes many membrane proteins. Supported by NIDCD grant R01 DC004295 to BS.
Large-Scale Proteomic Analysis of Cholesterol-Rich Membrane Microdomains in Cochlea
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Background
Compartmentalization of membrane proteins into microdomains is a key feature of cell signaling. These microdomains include buoyant, detergent-resistant membrane (DRM) specializations that are high in cholesterol and often enriched with cholesterol-binding proteins like caveolin and flotillin. Importantly, disruption of these domains with cholesterol-chelating cyclodextrins has proven extremely ototoxic (Ward et al., Pediatr Res, 2010). Using ultraperformance liquid chromatography-tandem mass spectrometry, we probed the protein composition of DRM fractions from the chick cochlea to gain insight into the structure and function of these domains.

Methods
Cochlear homogenates were pooled from 12 chicken basilar papillae (2 - 4 weeks old) and lysed in buffer containing 0.25 – 2% Triton X-100. Membrane proteins were separated by sucrose-gradient fractionation. Equal volume fractions were collected and probed for cholesterol content, protein concentration, and common markers of raft (caveolin) and non-raft (transferrin receptor) membrane domains. Twenty μg of protein from the DRM fractions were separated by SDS-PAGE, divided into ten equally sized gel segments, trypsin digested, and analyzed by nano-LC-MS/MS with an LTQ Orbitrap Velos Pro (MSBioworks).

Results
Homogenization in 0.5% Triton maximized protein concentration and enriched cholesterol in the DRM while optimally separating caveolin and transferrin receptor. From three independent samples, we identified 930 DRM proteins, with a false discovery rate of ~0.5%. The majority of these proteins were predicted by topology and ontology to be associated with cell membrane though only ~25% with the plasma membrane specifically. Identified proteins included caveolin, flotillin, prohibitin, erlin, and the GPI-anchored protein Thy-1, each common to lipid microdomains. Top terms in a process ontology analysis were metabolic processes (cluster frequency 35.4%), localization (24.4%), transport (21.3%), and cell communication (14.7%). Eight of the ten most abundant proteins in the DRM were associated with ion transport and energy metabolism.

Conclusion
Our results show that scaffolding proteins common to lipid rafts are enriched in the cochlea and that hundreds of other proteins segregate with these scaffolds. Cholesterol-rich microdomains have been widely associated with compartmentalized cell signaling, cell proliferation, excitability, exocytosis, and oxidative signaling. Our results revealed that proteins involved in energy metabolism and oxidative stress are abundant in cholesterol-enriched DRMs, offering a link between cholesterol disruption and hearing dysfunction which may explain cyclodextrin-induced ototoxicity.

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Generation of Atoh1-rtTA transgenic mice to transiently alter gene expression in hair cells

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Background
The Cre/loxP system has been commonly used for cell type specific control of gene expression, resulting in many significant advances for the hearing field. However Cre-mediated excision of the floxed region is permanent and in many cases, transient control of gene expression is desired. This can be achieved using the tetracycline-inducible system, where the reverse tetracycline transactivator (rtTA) protein binds to the promoter and activates transcription of a second transgene which contains a tetracycline response element (TRE). The rtTA protein can only bind to TRE and activate transcription in the presence of doxycycline, making it transient as the binding does not occur when doxycycline is removed. Cell-type specific expression can be achieved using cell-type specific promoters/enhancers to express the rtTA protein. Here, we used the well characterized Atoh1 enhancer to drive expression of rtTA. Atoh1 is a basic helix-loop-helix transcription factor that is required for the differentiation of hair cells in the inner ear and of granule cells in the cerebellum. In the inner ear, Atoh1 is expressed during late embryonic development and is down-regulated by postnatal day 6-7 (P6-7).

Methods
We used a construct that contains the Atoh1 enhancer followed by rtTA-IRES-hPLAP-SV40 Intron-poly A. The addition of the internal ribosome entry site (IRES)-human placental alkaline phosphatase (hPLAP) gene in the construct allows visualization of rtTA transgene expression (Chow et al., Dev. Dyn. 2006).

Results
We have obtained 12 transgenic founder mice and thus far 6 founders have given germline transmission. We are characterizing these transgenic mice using hPLAP staining in the inner ear and cerebellum and will further characterize their rtTA activity using TetO-lacZ reporter mice and doxycycline injection at early postnatal ages.

Conclusion
In addition to providing transient control of gene expression, these Atoh1-rtTA mice can be used in combination with other existing supporting cell-specific Cre or CreER mice for genetic manipulation of two genes in the inner ear in vivo. There is a second poster from our lab discussing a supporting cell specific rtTA allele (Sox10-rtTA; see Bradley Walters et al.).
Actin-Binding Protein 12 is a Component of the Hair Cell’s Cuticular Plate

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Background
The filamentous actin-based mesh of the hair cell cuticular plate is hypothesized to anchor the stereocilia in place at the apical surface of the cell. However, the precise role of the cuticular plate in hair bundle morphology and hearing remains unknown. Additionally, little is known about the molecular mechanisms that shape actin to build the unique structure of the cuticular plate.

Methods
By analysis of the zebrafish hair cell transcriptome, we identified several hair cell-expressed candidate molecules that could potentially be involved in hair bundle or cuticular plate morphogenesis based on their known actin-shaping activities in other tissues. We analyzed the localization patterns of these candidate molecules by immunolabeling in mouse hair cells as well as by expression of GFP-fusion proteins in zebrafish hair cells. We are also currently working to generate zebrafish mutants of each of these candidate genes using zinc-finger nucleases or transcription activator-like effector nucleases. Analyses of these mutants will help us to elucidate the role of these molecules in hair cell structure and hearing.

Results
We found that one candidate, Actin-Binding Protein 12 (ABP12), localizes throughout the cuticular plate in mouse vestibular hair cells. Ongoing analysis of GFP-ABP12 fusion protein expression in transgenic zebrafish hair cells also indicates a possible cuticular plate localization pattern.

Conclusion
We hypothesize that ABP12 is a key regulator of cuticular plate morphology and function. Future analyses of the effects of genetic perturbation of abp12 on hair bundle morphogenesis, function, and hearing will allow further elucidation of the molecular mechanisms shaping the cuticular plate and may also allow us to advance our understanding of the role of the cuticular plate in hearing.
Differential distribution pattern of the non-classical spectrin beta-V in the inner ear sensory hair cells during evolution

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Background
Within the mammalian auditory organ, outer hair cells (OHCs) amplify sounds by electromotility – the voltage-dependent contraction and elongation of the lateral plasma membrane (LPM). Electromotility is driven by the OHC transmembrane protein prestin. The actin- and spectrin-based cortical lattice that underlies the LPM is also important in this process; however, the precise composition of the cortical lattice, and the way in which it interacts with prestin (defective in a human deafness), have been unknown. In the mouse inner ear, we found that spectrin beta-V (which exists as an alpha-II/beta-V heterodimer) that is almost twice the length of conventional beta spectrins, forms the spectrin-based cytoskeleton that provides OHCs with the flexibility required for sound amplification.

Methods
Immunofluorescence microscopy analyses, Confocal laser scanning microscopy

Results
We detected spectrin beta-V in the vestibular epithelia where it displays a punctuated cytoplasmic distribution pattern. Interestingly, unlike in OHCs, no co-localisation was observed with alpha-II spectrin throughout the cytoplasm of vestibular hair cells. A co-localisation was, however, observed with Golgi-labeled structures, which suggests this spectrin involvement in protein/membrane trafficking. Because spectrin beta-V distribution pattern varies among inner ear sensory cell types, we analysed further its distribution pattern in Xenopus inner ear. We show that the protein is present also in the hair cells, essentially in the apical region of the cells, at the level of the cuticular plate. Faint labeling was also observed at the cell-cell junctions. This spectrin beta-V labeling in frog correlates with the distribution of the drosophila beta-Heavy spectrin observed in the apical region of epithelial cells.

Conclusion
Altogether, our findings support changes during evolution in the distribution pattern of the non-classical spectrin beta-V within the inner ear sensory hair cells.
Immunocytochemical localization of oncomodulin (Ocm) in the human inner ear

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Background
In the present study, we determined the distribution of oncomodulin (Ocm), a member of the parvalbumin family, relative to other EF-hand CaBPs by immunocytochemistry in the human inner ear using auditory and vestibular endorgans (macula utricle and cristae ampullaris) microdissected from human temporal bones obtained at autopsy and macula utricle obtained from ablative surgery. We also determined Ocm immunolocalization in the mouse cochlea and vestibule for comparative purposes (Simmons et al., J Comp Neurol, 518:3785-3802, 2010).

Methods
All human subjects (n=4, aged 82-85 years old) had documented normal auditory and vestibular function. The macula utricle acquired from patients undergoing surgery for acoustic neuroma (n=3, ages, 42, 44, 56 years old) was also used. Formalin fixed frozen sections obtained from the cochlea; macula utricle and crista were incubated with affinity purified rabbit polyclonal antibodies against recombinant full-length rat Ocm. Immunoreactivity was visualized using secondary antibodies labeled with Alexa 488 or horse-radish-peroxidase and DAB.

Results
In the human and mouse cochlea Ocm was exclusively located in outer hair cells (OHCs). The expression of Ocm was uniformly the base to apex. In contrast to the cochlea, Ocm expression was substantially reduced in human vestibular tissues at older adult ages (normal autopsy). In the human macula utricle from younger individuals (surgical specimens) Ocm immunoreactivity was located to hair cells in the central portion of the sensory epithelia resembling the striola.

Conclusion
The high degree of conservation of Ocm expression in the human and mouse suggest that Ocm may play an important role in calcium homeostasis. In addition it can be used as a marker for specific cells types in the vestibular sensory epithelia.
Slo is Targeted to the Basolateral Membrane Using a Dihydrophobic Amino Acid Motif
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Background
Large conductance K channels (BK) are important for determining the receptor potential of inner hair cells that allow a graded response to sustained sound induced depolarization and is essential for accurate sound encoding. These channels consist of an alpha subunit (Slo) and a variable number of auxiliary subunits. Slo is localized to the lateral wall of inner hair cells where they lie adjacent to ryanodine receptors, which are thought to be the site of Ca release and subsequent activation of these channels in a hair cells operating voltage range (-80 to -60mV). Similarly, these channels are thought to mediate efferent synaptic output in mid to high frequency outer hair cells. BK channels are localized at the basolateral surface of OHCs where its activation is thought to occur by Calcium induced Calcium release from the ER. The distribution of the ER in these OHCs mirrors that of efferent synaptic release sites.

Proteins are targeted to the basolateral surface of polarized epithelial cells using a number of intrinsic sequence motifs. In this study, we explore the role of a number of these motifs in targeting Slo to the basolateral surface of the cell.

Methods
We used MDCK cells as a model system and established that Slo is targeted to the basolateral surface of these cells. We identified four dihydrophobic motifs (DXXLL or D/EXXXLL) and one potential tyrosine motif (ΦXXY) that could target Slo to the basolateral surface of the cell. We then mutated these residues individually and transfected MDCK cells with Slo containing these mutations. The expression of these mutants on the basolateral surface was quantified using confocal microscopy.

Results
We determined that LI976 was critical for adequate targeting of Slo to the basolateral surface of MDCK cells. Substitution of these residues with either alanine or glutamine residues resulted in the loss of basolateral targeting with the protein retained within the Golgi. Finally, we establish that Slo uses a trafficking pathway that includes clathrin/AP1B since LLC-PK1 cells that lack the AP1B subunit show deficient Slo targeting to the basolateral surface of the cell.

Conclusion
Slo uses a dihydrophobic targeting sequence (LI976) and the clathrin/AP1B pathway to traffic to the basolateral surface of the cell.

cGMP Pathways in Mammalian Cochlear Hair Cells Underlying the Regulation of Intracellular [Ca\textsuperscript{2+}] by cGMP-gated CNGA3
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Background
The expression of olfactory/cone photoreceptor form(s) of cGMP-gated CNGA3 transcript in vestibular hair cells and CNGA3 protein in mammalian cochlear hair cells (Selvakumar et al., Biochem. J. Cell 443: 463-476, 2012) predicts a need to regulate cGMP pathways in hair cells. In vestibular hair cells, regulation occurs in part via cone photoreceptor CNG-phosphodiesterase 6c, PDE6c, but not PDE5.

Methods
Primers for the specific transmembrane isozymes of guanylyl cyclase and phosphodiesterase, and GCAP1 and GCAP2, that are utilized in vision and olfaction were applied in RT-PCR to rat organ of Corti and isolated cochlear hair cells. These primers crossed introns and all products were sequenced. Calcium fluorescence imaging was utilized to determine calcium responses in individually isolated cochlear hair cells to inhibitors of the cGMP-pathway isozymes.

Results
We show that transcript for cone photoreceptor PDE6c is expressed in singly-isolated inner and outer hair cells from the cochlea of the adult rat. Transcript for the cone photoreceptor guanylyl cyclase isoyme, GC-E, has been identified in cochlear inner hair cells. Transcript for photoreceptor isozyme GC-F, GCAP1 and GCAP2 have been found in an isolated organ of Corti (OC) cochlear subfraction. GCAP1 and GCAP2 regulate activation of GC-E and GC-F, and GCAP2 is recognized as being a protein binding partner of the hair cell ribbon synapse protein, ribeye. Transcript for the olfactory/photoreceptor form of CNGA3 has been identified in cochlear outer hair cells, and as functionally predicted for
this cGMP-gated ion channel, inhibitors of PDE6c caused an overall elevation of intracellular Ca\(^{2+}\) in singly-isolated outer hair cells over minute time intervals, as opposed to a decrease in intracellular Ca\(^{2+}\) that would be the consequence if voltage-gated calcium channel inhibition via PKG-phosphorylation were the primary determinant of intracellular Ca\(^{2+}\). Given the expression of cGMP-pathway proteins in the OC that are found in photoreceptors, we considered and confirmed transcript expression of photoreceptor rhodopsin and cone medium-wave sensitive opsin in rat organ of Corti, pointing to a mammalian correlate of the rhodopsin cascade identified in Johnston’s organ of Drosophila.

**Conclusion**
cGMP pathways in mammalian cochlear hair cells modulate hair cell intracellular Ca\(^{2+}\) and include components that are clearly identifiable as photoreceptor isozymes in retinal cGMP-cascades. These results support the suggestion that across evolution “different sensory modalities utilize common signaling cascades” (Senthilan et al., Cell 150: 1042-1054, 2012).

**USH1 proteins form an adhesion belt around the basolateral region of the photoreceptor outer segment, including the calyceal processes that are absent from mice**

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**Background**
The Usher syndrome is the first cause of inherited deaf-blindness in humans. USH1 Animal models wherein Usher syndrome type I (USH1) proteins — myosin VIIa (USH1B), harmonin (USH1C), cadherin-23 (USH1D), protocadherin-15 (USH1F) and sans (USH1G) — were targeted at different maturation stages of the auditory hair cells have proven efficient to unravel the role of USH1 proteins in proper organization of the hair bundle, the sound receptive structure, and functioning of the mechano-electrical machinery. By contrast, the mechanisms underlying USH1 retinal dystrophy remain unknown because mutant mice lacking any of the USH1 proteins do not display retinal degeneration. This discrepancy between the mouse and human retinal phenotypes has made it particularly difficult to elucidate the roles of USH1 proteins in the human retina. We addressed this issue here.

**Methods**
Immunohistochemistry, Confocal microscopy, Coimmunoprecipitation experiments, immunogold and scanning electron microscopy.

**Results**
We found evidence for the existence of major differences between rodents and primates in the structural and molecular architecture of photoreceptors at the interface of the inner and outer segments, and for a differential distribution of USH1 proteins associated with this particular region. In macaque photoreceptors, all USH1 proteins — myosin VIIa, harmonin, cadherin-23, protocadherin-15, sans — colocalized at membrane interfaces i) between the inner and outer segments in rods and ii) between the microvillus-like calyceal processes and the outer segment basolateral region in rods and cones. This pattern, conserved in humans and frogs, was mediated by the formation of an USH1 protein network, which was associated with the calyceal processes from the early embryonic stages of outer segment growth onwards. By contrast, mouse photoreceptors lacked calyceal processes and had no USH1 proteins at the inner-outer segment interface. Rodent photoreceptors also lack the long, rigid, F-actin roots of the calyceal processes, penetrating into the inner segment of the photoreceptor, and which confer a relationship between the inner and outer segments that is very different from that in photoreceptors lacking these structures.

**Conclusion**
Together, our findings accounts for why mice are poor models for retinal defects caused by USH1. They led us suggest that USH1 proteins form an adhesion belt around the basolateral region of the photoreceptor outer segment in humans, and that defects in this structure probably cause the retinal degeneration in USH1 patients.

The striated organelle and their spectrin connection in inner ear hair cells
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Background
Spectrins are a prominent family of cytoskeletal proteins initially described in red blood cells, where they were assigned the function of maintenance of the erythrocyte membrane's biconcave form. Non-erythrocytic forms of these proteins have since been identified, and their functional roles have grown just as steadily as the number of binding partners. In our work we are only beginning to appreciate the integral role that they play in inner ear hair cells in general, and the striate organelle in particular.

Methods
We used confocal and electron microscopy methods (Lysakowski et al., 2011), combined with co-immunoprecipitation (CoIP) and mass spectrometry analyses. For the CoIP experiments, each tissue type (vestibular, cochlear, and brain), was run in two parallel tubes - one for the bait antibody (mouse anti-α-II spectrin) and one for the control mouse IgG. The tissue extracts and sepharose G conjugated to antibody or IgG were incubated overnight at 40°C. After the reaction, the flow-through and CoIP elution products were analyzed using western blots and mass spectrometry for determining interacting protein identities. The mass spectrometry was done for us by our Core proteomics facility. Protein identifications were accepted if they could be established at greater than 95.0% probability and contained at least 2 identified peptides.

Results
We have evidence that the striated organelle is composed of αII and βII spectrin, whereas the other isoforms, viz. αI, βI, βIII, βIV, and βV spectrin, are not involved. αII and βII spectrin also localize to the cuticular plate and, hence, can be found in all hair cell types. On the other hand, the other non-striated organelle spectrins, although still present in hair cells, exhibit different or more restricted distribution patterns. Legendre et al. (2008, J. Cell Science 121: 3347-3356) recently showed that besides αII and βII spectrin, mammalian outer hair cells have β-V spectrin in their lateral walls, but they found no evidence of βI, βIII or βIV spectrins using immunofluorescence. By contrast, in our immunofluorescence experiments we have observed βI spectrin in outer hair cells but not in inner hair cells, and we are currently working out the distribution of βIII spectrin which preliminary evidence suggests is also present in cochlear hair cells.

Conclusion
We present further evidence of the distribution patterns of the spectrins and highlight how interacting partners differ, which might point to the different roles that spectrins play in inner ear hair cells.

Membrane recycling of outer hair cells
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Background
Efferent innervation of outer hair cells (OHCs) is probably important for modulating the mechanical impedance of the organ of Corti, and afferent innervation for feedback to the brainstem. Synaptic communication at the subnuclear pole of OHCs is thought to require tightly regulated exocytic and endocytic activities, membrane recycling. Rapid endocytic activity of OHCs in the guinea-pig cochlea has been already studied using the fluorescent membrane marker FM1-43 (Griesinger et al., 2004; Kaneko et al., 2006; Meyer et al., 2001). It was demonstrated that vesicles were intensively endocytosed at the apical pole of OHCs and trancytosed to different locations, namely into the basolateral membrane and through a central strand towards the nucleus. Since the significance of the endocytic activity in the subnuclear region of OHCs is still not clear, in this study we investigated rapid endocytic activity located at the synaptic pole of OHCs.

Methods
In response to application of FM1-43, confocal laser scanning microscopy was used to visualize membrane recycling of OHCs isolated from the functionally mature guinea-pig cochlea. Perfusion system was used that maintained constant laminar flow around the entire OHC. This system guarantees that upon application, the dye arrives at the same time and concentration to both the apical and basal poles of the OHC.

Results
After dye application, the fluorescence intensity immediately started to increase in both the infracuticular and subnuclear regions (t < 1 s). Signal intensity changes were recorded in the apical and basal poles relative to the signal at the
membrane. Measured 5 s after onset of FM1-43 application, the normalized signal intensity at depths of 1, 2, 4 and 6 µm deep beneath the membrane were 0.28 ± 0.02, 0.15 ± 0.03, 0.08 ± 0.03 and 0.06 ± 0.04 in the basal pole (N=5) and 0.39 ± 0.17, 0.16 ± 0.03, 0.09 ± 0.04 and 0.04 ± 0.03 in the subcuticular area (N=5), respectively. These data show no significant difference in fluorescence signal intensity changes between the opposite poles of the OHC - the subnuclear and the subcuticular areas.

Conclusion
Therefore, it is concluded that stained vesicles appearing in the subnuclear pole in that time interval cannot have derived from apical endocytic activity, but were locally endocytosed. Consequently, endocytic activities in both basal and apical poles of the OHC probably contribute equally to balance the exocytosis-evoked membrane surface increase - the membrane recycling.

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Prestin is Targeted to the Basolateral Membrane Using a Tyrosine Motif
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Background
The presence of the motor protein prestin in the basolateral wall of outer hair cells (OHCs) is critical for proper OHC function and cochlear amplification. Proteins are targeted to the basolateral surface of polarized epithelial cells using a number of intrinsic sequence motifs. In this study, we explored the role of a number of these motifs in targeting prestin to the basolateral surface of the cell.

Methods
We used MDCK cells as a model system and established that prestin is targeted to the basolateral surface of these cells. We identified five potential tyrosine motifs (ΦXXY) that could target prestin to the basolateral surface of the cell. We then mutated these residues individually and transfected MDCK cells with prestin containing these mutations. The expression of these mutants on the basolateral surface was quantified using confocal microscopy.

Results
We determined that Y520 was critical for adequate targeting of prestin to the basolateral surface of MDCK cells. We also established that substitution of Y520 with Phe and Ser resulted in a wild type phenotype and markedly disorganized targeting respectively. These data confirm that the signature important for basolateral targeting is dependent on the phenol ring and not on the OH side chain of the Tyr residue. Additionally, we confirm the lack of importance of an adjacent negatively charged group of residues in targeting prestin to the basolateral surface. Finally, we establish that prestin uses a pathway using clathrin/AP1B since LLC-PK1 cells that lack the AP1B subunit show deficient prestin targeting to the basolateral surface of the cell.

Conclusion
Prestin uses a tyrosine targeting sequence (520Y) and the clathrin/AP1B pathway to traffic to the basolateral surface of the cell.

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Can Oncomodulin Directly Regulate Prestin’s Function?
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Background
Prestin is a unique voltage–dependent motor protein responsible for the somatic electromotility of outer hair cells (OHCs) (Zheng et al., 2000), a process that is essential for providing the high sensitivity and frequency selectivity of mammalian hearing. Oncomodulin, also called β-parvalbumin, is a calcium-binding protein, selectively and abundantly expressed in OHCs (Sakaguchi et al., 1998). Oncomodulin’s mRNA developmental pattern is similar to prestin’s, starting with expression around P3, and peaking around P10 (Yang et al., 2004, Belyantseva et al., 2000). Furthermore, both prestin and oncomodulin mRNAs are down-regulated by the same microRNA (Lewis et al., 2009). It is also known that an exceptionally high level of oncomodulin protein is found in the soma of OHCs, which is surrounded by the prestin-embedded basolateral membrane (Hackney et al., 2005). In prestin-KO mice, augmented expression of oncomodulin is observed at both mRNA and protein levels (Miller et al., 2010). Similar expression patterns and regulation arrangements...
between prestin and oncomodulin indicate possible functional connections between these two important OHC-specific proteins.

**Methods**
In order to explore the association between prestin and oncomodulin, we investigated whether oncomodulin can directly bind to prestin. We created various prestin mutants by PCR. These mutant prestins include point mutations and deletions at various positions. Several biochemical methods including co-immunoprecipitation, SDS-PAGE/Western blot, and protein purification, were used to identify prestin’s amino acids that are important for oncomodulin binding.

**Results**
Experimental evidence suggests that prestin’s C-terminal domain has a significantly higher binding capability than control proteins.

**Conclusion**
Since prestin is expressed in the OHCs, whose mechanical property is significantly modulated by intracellular calcium (Dallos et al., 1997), it is reasonable to postulate that oncomodulin-prestin interaction at least partially underlies the calcium-dependent change in the mechanical properties of OHCs (Work supported by NIH Grants DC011813, DC010633, DC00089, and the Knowles Hearing Center).

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**Functional Consequences of Direct Calmodulin Binding to Prestin and other SLC26 Transporter C-terminal Domains: Inhibition of Transport by Calcium/Calmodulin**

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**Background**
Several studies have shown that calcium influences the function of various members of the SLC26 protein family (Lamprecht et al., 2009; Loriol et al., 2008), including prestin (Frolenkov et al., 2000), the central force-delivering protein in the mammalian cochlear amplifier (Zheng et al., 2000). The mechanistic connection between calcium and SLC26 proteins, however, has heretofore not been elucidated. We have previously presented bioinformatic and biochemical evidence that calmodulin binds directly to intrinsically-disordered regions (IDRs) in the intracellular carboxy-terminal domains of several, if not all, SLC26 members in a calcium-dependant manner (Jacob and Dallos, 2009).

**Methods**
We performed ratiometric measurements of intracellular pH changes in cultured cells expressing various SLC26 proteins, the kinetics of which reflect the magnitudes of anion transporter activities of the expressed transporter proteins. The effects of the calcium ionophore, calcimycin, and the calmodulin inhibitor, trifluoperazine (Vandonselaar et al., 1994), were examined in order to demonstrate functional ramifications of calmodulin binding to SLC26 proteins.

**Results**
Calmodulin binding to IDRs within intracellular domains of SLC26 proteins is expected to be induced by increased intracellular calcium upon calcimycin treatment. Our experiments indicate that this binding significantly reduces the kinetics of pH change, and that the inhibition of calmodulin by trifluoperazine significantly alleviates this calcimycin-induced inhibitory effect.

**Conclusion**
Our results strongly indicate that the functional consequence of calmodulin binding to the IDRs in the intracellular carboxy-terminal domains of SLC26 transporters is inhibitory. We also discuss some ideas about potential ramifications for prestin electromotility and cochlear amplification.

This work is supported by NIH grants DC00089 (to MAC), DC010633 (to JZ), and by the Knowles Hearing Center.
Effects of the Elevation of Intracellular Calcium on the Outer Hair Cell Plasma Membrane Motors

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Background
It has been proposed that calcium signaling in outer hair cells (OHCs) may regulate cochlear function. An increase of intracellular calcium is known to affect the stiffness of the cortical cytoskeleton in OHCs. It has been also proposed but never directly demonstrated that the increase of intracellular calcium may regulate the operation of the prestin-based plasma membrane motors. Calcium concentration within an OHC may be elevated either through the calcium influx from outside of the cell or through calcium release from the subsurface cisternae, an endoplasmic reticulum-like structure of the OHCs. The OHC cortical cytoskeleton and the membrane motors elements are located at the lateral wall of the OHC in close proximity to the subsurface cisternae.

Methods
We used photo-activatable (caged) calcium chelators to release free calcium within the OHC and fast calcium imaging to monitor the propagation of calcium signal along the cell. Whole cell patch clamp recordings were used to simultaneously measure the OHC non-linear capacitance.

Results
In acutely isolated organ of Corti explants loaded with o-nitrophenyl EGTA (NP-EGTA) UV laser illumination with duration less than 20 ms produced rapid rise in intracellular calcium concentration measured by calcium indicator fluo-4. After an initial calcium spike we observed a diffusive propagation of calcium longitudinally along the OHC with an estimated speed of ~ 1 μm/s. The calcium-induced calcium release, if present, was difficult to resolve. Puff application of 5 μM ryanodine to OHCs resulted in only moderate calcium responses. Thapsigargin (50 μM), an inhibitor of the endo/sarco-plasmic reticulum ATPase, does not apparently affect the propagation of uncaged calcium along the OHC. In the presence of cesium in the patch pipette, the whole-cell currents were not affected by the release of calcium into the cytosol. However, the non-linear capacitance decreased by ~10% after about one minute after surge of intracellular calcium.

Conclusion
1) Our data suggests a minimal contribution of calcium-induced calcium release in propagation of calcium signals along OHC. 2) However, a decrease of non-linear capacitance after calcium uncaging indicates potential modulatory effect of intracellular calcium on prestin-based plasma membrane motors.
The relationship of prepulse effect and prestin density in an inducible prestin cell line
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Background
Mechanical activity of the outer hair cell (OHC) underlies the enhanced sensitivity and frequency resolving power of the mammalian cochlea. Prestin, a member of the SLC26 anion transporter family, has been identified as the molecular motor for electromotility. The voltage dependence of prestin’s nonlinear capacitance (NLC) depends on long term holding potential — we call this a prepulse effect. The relationship of prepulse effect and prestin density in an inducible prestin cell line remains to be elucidated.

Methods
HEK293 cells were cultured in DMEM containing 50 U/ml each of penicillin and streptomycin, 10% fetal bovine serum at 37°C in a CO2 incubator (5%). All whole cell recordings were made on single cells growing on a coverslip at 2h, 4h, 6h, 10h and 24h after tetracycline induction. The NLC traces show the change of membrane capacitance Cm in response to voltage across the membrane Vm, which was fit with a Boltzmann equation.

Results
We assayed four different parameters of NLC, Qsp, z, Vh and ΔVh, to delineate prestin function. Qsp, charge movement per unit of membrane surface, increases with time in a sigmoidal manner. In contrast to Qsp, which takes days to reach its asymptotic level, the z value reaches a maximum level of 0.8e 6h after induction. Vh values reach its asymptotic value 6h after induction, shifting in the depolarizing direction with time. ΔVh, the difference between Vh with long term holding potentials (1 min) of -100mV and +50mV holding potentials, an index of the prepulse effect upon Vh, increases steadily from 2h and reaches its peaks 10h after induction.

Conclusion
The prepulse effect upon Vh increases with prestin density, and may correspond to early molecular maturation events.
Morphological and functional changes in a new animal model for Meniere's disease
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Background
Meniere's disease (MD) is histologically characterized by endolymphatic hydrops (EH) in the inner ear. There is considerable evidence that water homeostasis in the inner ear is regulated partly via the vasopressin-aquaporin2 (VP-AQP2) system. EH is considered to be the result of incoordination of inner ear water homeostasis, which involves excessive production of endolymph and/or reduced absorption of endolymph. Until recently, however, the detailed mechanisms underlying the over-accumulation of endolymph were unclear. We aimed to examine both morphological and functional changes of the inner ear system in our novel combined animal MD model, involving the combination of surgical obliteration of the endolymphatic sac and duct and the administration of desmopressin (vasopressin type 2 receptor agonist; V2 agonist).

Methods
In experiment No. 1, guinea pigs used in morphological studies underwent surgical obliteration of the endolymphatic sac and duct in one ear and were maintained for a week or 4 weeks, and divided into with or without desmopressin administration. We quantitatively assessed EH in the cochlea, vestibules and semicircular canal. In experiment No.2, we performed vestibular examination. We recorded spontaneous nystagmus and observed the presence or absence of balance disorder in normal control group, desmopressin administration group and surgical treated groups with or without desmopressin administration.

Results
In experiment No. 1, both the increase ratio (IR) of the scala media area and the proportion of the endolymphatic space in the saccule were significantly higher in the Combined group than in the Surgical groups. There were no significant differences in the degree of hydrops in the utricle or semicircular canal among ear groups. In experiment No. 2, all animals of the Combined groups showed spontaneous nystagmus and balance disorder; meanwhile, the Surgical groups were found to be asymptomatic.

Conclusion
Our experimental animals not only presented severe EH in the cochleae and saccules, but also showed episodes of balance abnormalities along with nystagmus. EH may be exacerbated due to endolymphatic sac dysfunction combined with the effect of desmopressin and acute V2 effect, which may accompany vestibular abnormalities that are similar to the vertigo attack in patients with MD.
Do autoimmunity to IgG1 and their immune complex reaction cause Meniere’s disease?
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Background
This study was performed to identify autoimmune reaction as a main pathologic mechanism of Meniere’s disease. Authors tried to identify main target antigens in Meniere’s disease which causes inner ear damage and endolymphatic hydrops of Meniere’s disease.

Methods
First, proteins which only exist in the endolymphatic sac (ES) luminal fluid of Meniere’s disease were identified by proteomic analysis (LC-MS/MS, n=3). Based on the result, authors conjectured pathophysiology of Meniere’s disease and performed protoarray with the sera of Meniere’s disease patient (n=10) and compared the result with those of normal controls (n=10). With the data of protoarray, we identified target antigens reacted with autoantibodies of patients’ sera. Then, we examined if the target antigens also exist in the human endolymphatic sac tissues and ES luminal fluid of patients using proteomic analysis (LC-MS/MS).

Results
Proteins only exist in the luminal fluid of Meniere’s disease but not in the control were 9 proteins; eight of them were immunoglobulin and one was interferon regulatory factor. This infers that immunologic or autoimmune reaction could be the pathologic mechanism of Meniere’s disease. In the protoarray, 18 antigens showed increased reaction more than two folds when compared with normal controls. Among the antigens, only one protein, IgG1, exist in the human ES sample and ES luminal fluid and it showed increased reaction with the sera of Meniere’s disease more than 200 folds.

Conclusion
This result means that autoimmune reaction to IgG1 and immune complex formation can be the pathologic mechanism of Meniere’s disease like other rheumatologic diseases such as rheumatoid arthritis and SLE. For validating the results, immunoprecipitation with the ES luminal fluid and IgG1 protein and mass analysis of immune reaction with Meniere’s disease patients’ sera should be performed.
An Electromotility Mechanism Driven by the Tectorial Membrane
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Background
The mammalian cochlea is a remarkable sensor capable of detecting and analyzing sounds that generate subatomic vibrations. This extraordinary sensing property is widely believed to depend on electromotility of outer hair cells. Here we describe an additional electromotility mechanism driven by the tectorial membrane (TM), whereby transduction currents acting on TM fixed charge groups generate local TM deformations that could contribute to hair bundle motion.

Methods
We developed a microaperture chamber to test TM electrical properties by positioning isolated mouse TMs as an electrochemical barrier between two fluid-filled compartments. This setup allowed both measurement of TM interfacial voltages and application of electrical stimuli.

To determine TM fixed charge density, we measured DC interfacial potentials between the TM and the surrounding baths as a function of bath ionic concentration. The resulting voltages were fit by a model based on the Donnan relation.

To measure TM electromotility, AC electric fields were applied to the TM in its middle and marginal zones, which normally overlie the hair bundles. Electrically-evoked displacements in the bulk of the TM were visualized and quantified using a stroboscopic computer vision system and Doppler optical coherence microscopy.

Results
The net charge of the TM was large (~7.1 mmol/L at physiological pH), representing 1 fixed charge molecule for every 25 cations in endolymph. The high density of fixed charge contributes significantly to TM compressive stiffness and also suggests that electrical stimuli could generate an electromechanical response.

Electrically-evoked TM motions were nanometer-scaled (~5-200 nm), increased linearly with electric field amplitude (0.05-20 kV/m) and decreased with frequency (1-1000 Hz). This frequency dependence can be understood in terms of the interplay between electrophoresis and electro-osmosis.

Conclusion
Results reported here show that the TM contains a high density of fixed charge that contributes to its compressive stiffness and can generate electrically-evoked displacements. These displacements may have important implications for cochlear mechanisms. Although the electric fields applied in this study were large, they are in the range that can be generated near hair cell transduction channels. The low pass nature of the TM's frequency dependence suggests that this mechanism may be particularly relevant at low frequencies and in the cochlear apex. TM electromotility thus could be important in the deflection of cochlear hair bundles in vivo.
Hearing Impairment at Low and High Frequencies after Loss of Mammal-specific Tectorial Membrane Component Carcinoembryonic Antigen Cell Adhesion Molecule 16 (CEACAM16)

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Background
The mammal-specific, secreted CEACAM16 is a member of the carcinoembryonic antigen gene family. CEACAM16 is exceptionally well conserved among species. CEACAM16 is selectively expressed in the inner ear. We studied the hearing function and the cochlear expression of CEACAM16 to elucidate its importance for auditory function in young and aged Ceacam16−/− mice.

Methods
Ceacam16−/− mice were analysed for age related and trauma induced hearing deficits by ABR and DPOAE measurements and exposure to noise. In cochlear sections of Ceacam16−/− mice the tectorial membrane was inspected and gene expression was studied by in situ hybridisation and immunohistochemistry. Protein expression was furthermore studied by Western blot analyses.

Results
CEACAM16 is specifically expressed in the inner ear. The hearing phenotype of Ceacam16−/− mice was similar to the observed hearing impairment in members of human Family 1070 with non-syndromic autosomal dominant hearing loss (DFNA4) who carry a missense mutation in CEACAM16.

Conclusion
CEACAM16 can probably form higher order structures with other tectorial membrane proteins and influences the physical properties of the tectorial membrane. Evolution of CEACAM16 might have been an important step for the specialization of the mammalian cochlea, allowing hearing over an extended frequency range.

Remodeling and Biosynthesis of Glycosaminoglycans Revealed Using a BODIPY-Xyloside Conjugate in the Vestibular System

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Background
Examination of proteoglycan biosynthesis and remodeling in the semicircular canals and utricle using a novel BODIPY-xyloside conjugate. Xylosides are known to enter the Golgi apparatus and prime the production of glycosaminoglycans (GAGs). When conjugated with a fluorescent reporter and introduced into the inner ear endolymph, we found newly expressed fluorescent GAGs in vivo. We provide a new powerful tool to study inner-ear glycobiology and associated disorders.

Methods
Adult oyster toadfish (Opsanus tau) of either sex weighing 500 gm with rostral-caudal length greater than 25 cm were obtain from the Marine Biological Laboratory (Woods Hole, MA), and all experiments were carried out in accordance to the approved protocols at the Marine Biological Laboratory and the University of Utah. The BODIPY conjugated xyloside used in this study (BX, VY-IIB 57A) was synthesized following a previously described technique (Kuberan et al., ChemBioChem. 2008). Two microliters of a 2 μM BX in DMSO is diluted in endolymph and administered via glass micropipette into the dorsal extreme of the anterior endolymphatic canal. Single unit afferent recordings and fluorescence imaging were done in vivo. After ~4 hours, the inner ear was harvested and fixed for either confocal or 2-photon imaging (Olympus FV1000/MPE), or biochemical analysis. Western blotting was performed using mouse anti-heparin sulfate proteoglycan (AbDSerotec), and mouse anti-chondroitin sulfate proteoglycan (Invitrogen).

Results
Within minutes BX post injection, fluorescence was detectable in the crista of the anterior and horizontal canals. BX was not fluorescent in the lumen of the canal or ampulla, showing incorporation into the cells was a precursor to fluorescence. The time course and spatial distribution of fluorescence development suggests ongoing GAG production occurring at different rates among different vestibular structures. Hair cells expressed strong and persistent fluorescence, with the brightest fluorescence localizing at the apical end of the cell. Multi photon imaging showed strong phalloidin labeling of stereocilia and a BX primed fluorescent kinociliumglycocalyx. The epithelial lining of the endolymphatic space and
supporting cells also expressed fluorescent GAGs. BX revealed GAG ultrastructural features of the cupula including long fluorescent projections from the crista to the ampullary apex.

**Conclusion**
A novel xyloside conjugate administered into the endolymph, demonstrates priming of fluorescent GAGs in the inner ear. Additionally, we report strong expression in the vestibular organs of chondroitin sulfate and heparin sulfate proteoglycans. This study reveals GAG ultrastructure, time course of GAG production, remodeling and repair in the ear.

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**Experimental OAE growth rates and general properties of a dynamical system**

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**Background**
The dynamical properties of the active feedback mechanism localized in outer hair cells can be recognized in the otoacoustic emissions (OAE) I/O curves. OAEs seem therefore to be able to reflect the fundamental characteristic of the cochlear amplifier.

**Methods**
High-resolution SFOAE and DPOAE spectra have been measured at different stimulus levels in young healthy subjects using a chirp technique. TEOAE were measured with the double evoked paradigm. A time-frequency filtering method was applied to separate OAE components from different generation mechanisms according to their different phase-gradient delay. In the TEOAE and SFOAE cases, time-frequency analysis permitted separation of the main reflection component from a shorter latency component, and from multiple-reflection components. In the DPOAE case, the overlap component and the main reflection component have been separated.

**Results**
SFOAEs are thought to be generated by a place-fixed mechanism due to linear reflection from cochlear roughness. The SFOAE complex response is consistent with that measured in the time domain for TEOAEs. If the peak BM response is compressive as \( p^{1/3} \), a saturated response \( p^{1/3} \) is expected in the case of peak reflection both in the TEOAE and in the SFOAE. This prediction is well confirmed by the present work. A reduced compression level is shown by the TEOAEs and SFOAEs shorter latency components, which seem therefore attributable to reflections from more basal cochlear regions.

A linear growth rate has been found over a wide stimulus level range for the DP overlap component. This finding is coherent with a cubic distortion source in which the BM displacement follow a compressive behavior as \( p^{1/3} \). As the overlap component is the source for the reflection component, when the first one grows linearly, the reflection one is expected to grow as \( p^{1/3} \). This prediction is also confirmed by the experimental results.

**Conclusion**
The characteristics of the growth rate of the time-frequency separated OAEs components seem to reflect the dynamics of the cochlear active gain.
A compressive BM response growing as \( p^{1/3} \), as predicted for a dynamical system operating at the Hopf bifurcation, is consistent with the experimental data.
What You See is Not What You Get: DPOAEs versus Intracochlear DPs
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Background
The characteristics of distortion product otoacoustic emissions (DPOAEs), ie, distortion products (DPs) measured in the ear canal have been thoroughly described. In contrast, there is relatively little known concerning the behavior of intracochlear DPs (iDPs) propagated toward the DP frequency place (f_{dp}) on the basilar membrane (BM). Detailed comparisons of iDPs to DPOAEs could provide valuable insights on the extent to which DPOAEs can be assumed to mirror DPs within the cochlea.

Methods
Whitehead et al (1995) described a technique whereby the behavior of iDPs could be inferred by interacting a probe tone with the DP of interest to produce a secondary DPOAE. The behavior of the 'secondary' DPOAE was then used to deduce the characteristics of the iDP. In the present study, this technique was used in rabbits to directly compare DPOAE f_2/f_1-ratio functions, level/phase maps, and interference response areas (IRAs) to their iDP counterparts. 2f_1-f_2 and 2f_2-f_1 DPOAEs were collected with f_1 and f_2 primary-tone levels from 75-35 dB SPL. During the collection of these DPOAEs, a 50-dB SPL f_3 was placed at a DP/f_3 ratio of 1.25 to evoke a secondary DPOAE at 2f_3-(2f_1-f_2) or 2f_3-(2f_2-f_1). In some experiments, a fixed 4th interference tone (IT) was presented, or was alternatively swept in frequency and level to produce an IRA describing the effects of the IT on DPOAEs and iDPs.

Results
It was found that low primary-level DPOAE f_2/f_1-ratio functions peaked around f_2/f_1=1.25, while the corresponding iDP ratio functions peaked at f_2/f_1≈1.0. Also, notches observed in DPOAE f_2/f_1 ratio functions were not present in their iDP counterparts. Additionally, ITs above f_2 at narrow f_2/f_1 settings often enhanced the DPOAE, while the iDP remained unaffected. And, level/phase maps showed rapid phase variation with f_{dp} for the narrow ratio 2f_1-f_2 and all 2f_2-f_1 DPOAEs, while the corresponding iDPs evidenced relatively constant phase.

Conclusion
DPOAEs often provide poor representations of iDPs, because distributed DPOAE components directed toward the ear canal are often in cancellation, while iDP components directed toward f_{dp} add in phase.

Mechanical two-tone-distortions in the movement of the tympanal membrane of insects
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Background
During stimulation with two pure-tones, tympanal organs of insects emit pronounced distortion-product otoacoustic emissions (DPOAEs) that are indicative of non-linear ear mechanics. There is evidence that the scolopidia, the auditory mechanoreceptors in tympanal organs, are involved in their frequency-specific generation [Möckel et al.: J Comp Physiol A193, 2007]. Our current study aims to measure the mechanical correlates of DPOAEs within the movement of the tympanal membrane of the locust Schistocerca gregaria.

Methods
We analysed the movement of the tympanal membrane during stimulation with two pure-tones f_1 and f_2. Both stimuli f_1 and f_2 had been optimized to evoke large emissions using a closed system to record sensitive DPOAEs in insects. The same frequencies were then used for the open system measurements required by the laser Doppler vibrometry. The laser measurement points were arranged around the potential generation site of the emissions, that is, the attachment point of the scolopidial mechanoreceptors at the tympanal membrane. Stimulation levels of f_1 and f_2 started at 30 dB SPL and reached up to 70 dB SPL.

Results
During stimulation with two pure tones f_1 and f_2, the amplitude response of the tympanal membrane showed additional peaks at the emission frequency 2f_1-f_2. The 2f_1-f_2 mechanical displacement level was 43 dB below the f_2 amplitude which is comparable to the OAE recording situation when using a closed acoustical measurement system. The 2f_1-f_2 tympanum displacements amounted up to about 0.25 nm for single measurement points located close to the attachment
area of the scolopidia. The threshold to evoke a 2f1-f2 mechanical response in the tympanum movement during these open-system measurements lay at about 50 dB SPL stimulation level.

Conclusion
We demonstrate the existence of mechanical two-tone-distortions in the movement of the tympanal membrane at the 2f1-f2 frequency. The largest displacement amplitudes and therefore potential place of their generation lies close to the attachment point of the dendrites of the receptor cells.

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Time-frequency analysis of transiently evoked otoacoustic emissions measured with different acquisition protocols
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Background
Transiently evoked otoacoustic emissions (TEOAEs) are commonly recorded as average responses to a repetitive click stimulus. If the click train has constant polarity, a linear average results; if it contains a sequence of clicks of differing polarity and amplitude, a nonlinear average can be calculated. The purpose of this study was to record both protocols from the same set of ears and characterize the differences between them.

Methods
Time-frequency (TF) analysis of the recorded signals was done by decomposing them into their basic waveforms. For this, a method of high-resolution adaptive approximation was used, a technique based on the matching pursuit (MP) algorithm.

Results
It was shown that the main difference between TEOAEs recorded with the linear and nonlinear methods is found in the 0-5 ms post-stimulus time window. It was assumed that the signal derived from the linear protocol was contaminated by stimulus artifact. After removal of this artifact, the time-frequency properties of TEOAEs recorded under both the linear and nonlinear protocols were similar. However reproducibility and signal to noise ratio were higher for the linear TEOAEs. Additionally, it was shown that TEOAEs recorded under the linear protocol appear to be less dependent on the presence of synchronized spontaneous otoacoustic emissions (SSOAEs).

Conclusion
TF analysis has shown that the main difference between CEOAEs recorded with the linear and nonlinear methods is found in the 2.5-5 ms post-stimulus time window, and it affects frequencies below 2.2 kHz. The difference is mainly due to stimulus artifact in the linear protocol. This artifact can be removed, either by the MP method or by a combination of windowing and linear filtering, a process that still leaves the high-frequency, early OAE response intact. TF analysis verifies the reliability of the proposed method.

The Effect of Probe Frequency on the Time-Course of Recovery from the Medial Olivocochlear Reflex
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Background
Different methods for estimating the effect of the medial olivocochlear reflex (MOCR) on stimulus-frequency otoacoustic emissions (SFOAE) in humans appear to yield different estimates of the time-course of recovery from the effect. The effect of the MOCR elicited by broadband noise, calculated by extracting changes in the relationship between the earcanal pressure produced by two brief simultaneous probe tones relative to the sum of the pressures produced by the tones presented in separation decayed over ~600-700 ms after the offset of the elicitor [K. P. Walsh, E. G. Pasanen, and D. McFadden (2010). Overshoot measured physiologically and psychophysically in the same human ears. Hear. Res. 268, 22-37]. In comparison, the MOCR effect elicited by a notched-noise, estimated from changes in the ear-canal pressure using a continuous fixed-level probe was substantially shorter, on average ~160 ms [B. C. Backus and J.J. Guinan, Jr. (2006). Time-course of the human medial olivocochlear reflex. J. Acoust. Soc. Am. 119, 2889-2904]. Aside
from using different methods for estimating effects of efferent activation, the two studies also used different probe frequencies (4 versus 1 kHz). Thus the question remains as to whether the differences in recovery times are due to differences in methodology or due to a frequency dependence of the recovery from the MOCR.

**Methods**
The time-course of recovery was estimated using continuous fixed-level (40-dB SPL) probes with frequencies of 1, 2, 4 and 6 kHz. For each probe, a 60-dB SPL ipsilateral broadband noise with a 2-octave notch around the probe frequency was used to elicit the MOCR. The ear-canal pressure was measured in the presence and absence of the elicitor to extract changes in SFOAE at the probe frequency due to the elicitor. A heterodyne technique was used to calculate complex-valued ear-canal pressure at the probe frequency.

**Results**
Changes in SFOAE following the offset of the elicitor, expressed in terms of the proportion of the magnitude of SFOAE measured using a suppression paradigm for each probe frequency, were fitted with an exponential to estimate the time of recovery. The time constants from the best exponential fits to the normalized changes in SFOAE magnitude increased with increasing probe frequency.

**Conclusion**
The data suggest that the time of recovery from the effect of the MOCR estimated using SFOAEs increases with increasing probe frequency.

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**Probing Otoacoustic Emissions in the Budgerigar (Melopsittacus undulatus)**

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**Background**
Given the complex nature of their vocalizations, birds have become an increasingly attractive model for studying neurophysiological mechanisms underlying the processing of complex sounds. However, much is still unknown about the function of the peripheral auditory system in birds, and there are significant physiological differences between birds and mammals (e.g., hair cell distribution, lower prestin density, structure of the tectorial membrane). The present study explores this issue by characterizing several types of otoacoustic emissions (OAE) in the budgerigar (Melopsittacus undulatus).

**Methods**
Birds were lightly anesthetized using either isoflurane (gas) or ketamine and xylazine (injectable, sub-cutaneous) and were expected to fully recover following the recording session (typically 1-2 h long). Feathers about the external auditory meatus were gently trimmed and the OAE probe was then sealed to the head using vaseline.

**Results**
While no spontaneous emissions were observed, physiologically-sensitive evoked emissions were readily measurable. Though not thoroughly explored, emissions magnitudes exhibited a degree of sensitivity upon both the depth and type of anesthesia used, consistent with previous reports.

**Conclusion**
OAEs were confined to the most sensitive regions of the audiogram. Tuning estimates derived from stimulus-frequency OAEs suggest auditory filter bandwidths similar to those of chicken (Gallus gallus domesticus). Relative to mammals, distortion-product OAEs exhibited both similarities (e.g., shallow 2f1-f2 DPOAE phase-gradients for larger fixed f2/f1 ratios) and differences (e.g., reduced magnitude fine structure).

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Comparison of MOC efferent effects on SFOAEs, DPOAEs and compound action potentials.

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Background
Otoacoustic emissions (OAEs) have been used as a non-invasive way to assess the effects of medial olivocochlear (MOC) efferents in humans and laboratory animals. However, the MOC effect that is relevant for hearing is the inhibition of auditory-nerve (AN) fiber responses. Nonetheless, there has been little work on the relationship between MOC effects on OAEs and neural responses.

Methods
We measured stimulus-frequency OAEs (SFOAEs), distortion-product OAEs (DPOAEs) and AN compound action potentials (CAPs) evoked by tone pips in deeply anesthetized albino guinea pigs. MOC efferent neurons were activated by electrical shocks at the floor of 4th ventricle. DPOAE tones had a frequency ratio of 1.2 with the higher-frequency tone 10 dB SPL above the lower-frequency tone. SFOAEs were separated from the probe sound by taking the difference of the ear-canal sound at the SFOAE frequency with and without a suppressor tone 50 Hz above the probe frequency and 20 dB SPL above the probe level. SFOAE, DPOAE and N1 level functions were measured with and without MOC stimulation. To keep the MOC effects frequency specific, probe tones above 50 dB SPL were not used. The MOC effects for SFOAEs, DPOAEs and CAPs were quantified by the MOC-induced shift to higher levels of the amplitude-level functions. Comparisons of MOC effects on CAP and emissions were done at 5-10 dB above the CAP threshold level. For the vast majority of collected data, this corresponded to 25-35 dB SPL.

Results
Our preliminary results suggest that in animals with good hearing the MOC inhibition of SFOAEs and CAPs are comparable for tones near 7-8 kHz. For tones below 7-8 kHz, the MOC inhibition tended to be larger on CAPs than on SFOAEs. In several animals there were tone frequencies for which the SFOAE amplitude-level and/or phase-level functions were non-monotonic with MOC stimulation and monotonic without MOC stimulation. Preliminary results suggest that the MOC effects on DPOAEs are smaller than MOC effect on SFOAEs and CAPs.

Conclusion
In the mid-frequency region, MOC effects on SFOAEs and on gross neural responses are similar. More work is needed to characterize the relationship outside of this frequency region, and between MOC effect on DPOAEs and CAPs in all frequency regions. Supported by RO1DC00235, P30DC005209. Maria Berezina-Greene is also supported by NSF GRFP.

Determination of Primary- and Secondary-Source Components Using Short-Pulse Distortion Product Otoacoustic Emissions

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Background
According to a widely-used model, distortion product otoacoustic emissions (DPOAEs) evolve from interference of a so-called primary- and secondary-source component within the cochlea. The presence of two components can be visualized during the onset of the f2 primary if signal blocks recorded with suitable primary-phase variations are averaged in the time domain (Whitehead et al., 1996). Recently, we showed that sampling the resulting time signal at an appropriate time instant leads to reliable extraction of the primary-source component (Vetesnik et al., 2009). However, the secondary-source component is not always reliably quantified, especially if both contributions are approximately in phase. Here, we present a new method using short-pulse DPOAEs that enables the quantification of both sources independent of their respective phase.

Methods
DPOAEs were acquired from four subjects with distinctive fine structure. The f2 tone was pulsed for 8ms, whereas the f1 tone was presented continuously. The ratio f2/f1 was kept constant at 1.2, with f2 increasing from 1.7kHz to 2kHz in steps of 20Hz. Primary-tone levels were between 25 and 65 dB SPL. For the investigated frequency range, the primary-source component attains its steady state approximately 8–10ms after onset of the f2 tone, while the secondary-source component is delayed by another 8–10ms. Presenting the f2 primary with a short pulse of 8ms leads to suppression of the primary-source component during the rise of the secondary-source component. Cancellation of the primary tones and
other DPOAE waveforms, except for the cubic component of interest at 2f1–f2, was achieved using ensemble averaging with suitable phase shifts between adjacent signal blocks (Whitehead et al., 1996).

Results
DPOAE amplitude and phase responses show considerable differences depending on the relative interference phase, particularly at time instants shortly after the decay of the primary-source component. DPOAE signals with destructive interference show a distinctive notch separating the contributions of the two components and rapid phase jumps close to 180°. A phase variation of 90° indicates the quadrature condition, whereas an invariant phase implies constructive interference. In the latter case, using the short-pulse technique, the size of the secondary source can be detected 16–20ms post-onset.

Conclusion
Short-pulse DPOAEs present a promising approach to separate and quantify the two source components with a single-frequency measurement, thus avoiding the otherwise necessary acquisition of DPOAEs over a broader frequency range.

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Insights from a structural theory of pitch: Existence region, sound level dependence and discriminability.
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Background
Pitch is a fundamental auditory percept which is perceived by many species and for humans constitutes the basis of melody production. Clearly, what distinguishes sounds with a pitch from other sounds is the temporal regularity of the signal. In other words, sounds with similar period are perceived with similar pitch. However, solely defining pitch as the perceptual correlate of periodicity raises a few discrepancies. Periodic sounds elicit pitch only within a certain domain of existence. There are small but significant level-dependencies of pitch for low frequency pure tones. Finally, although independent of the exact harmonic content (e.g. missing fundamental); the resolvability strongly determines the salience and discriminability of pitch. Here, we propose that pitch is the perceptual correlate of the similarity structure in the vibration pattern of the basilar membrane.

Methods
We use both analytical method and data analysis to study the implications behind the detection of the similarity structure. Analytical derivations are used to obtain an implicit equation linking the phase difference at two basilar membrane locations, the sound level and the estimated pitch. We analyze previously published guinea-pig auditory nerve fibers (ANF) responses to low frequency tones (Palmer and Shackelton 2009)and fit von Mises distribution to the period histogram to obtain an analytical description of the phase probability distribution.

Results
We describe the similarity structure along the basilar membrane by identifying positions whose vibrations are equal but with possible time delay. From this mathematical definition, we derive two types of structure: across channel and within channel similarity. We show that across-channel similarity is only possible for resolved harmonics while within channel similarity is possible for both resolved and unresolved harmonics. In our framework, the maximal delay allowed set the lower limit of perceived pitch and we demonstrate the implications of resolvability on this lower limit. In the case of a pure tone, the pitch shift with level is computed using phase distribution from ANF and compared to results from the psychophysics. Finally, we estimate tone discriminability as a function of F0 using vector strength.

Conclusion
Our work provides new insights into possible mechanisms used for pitch perception.

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Effect of Efferent Activation on Cochlear Gain and Compression Estimates
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Background
Behavioural and physiological evidence suggests that the amount of gain applied to the basilar membrane may change during the course of acoustic stimulation due to efferent activation of the cochlea (Liberman, 1996). This study used the
Fixed Duration Masking Curve (FDMC) method (Yasin et al, 2012) to estimate human cochlear gain and compression, both with and without a precursor sound. In the FDMC method, masker level at threshold for a 10-dB SL, 4-kHz signal is obtained in the presence of an on-frequency (4 kHz) and off-frequency (1.8 kHz) forward masker, as a function of signal duration, with total masker-and-signal duration set to 25 ms, and masker-signal silent interval set to 0 ms. A short-duration masker ensures that accurate estimates of basilar membrane gain and compression can be obtained in the absence of efferent activation. Presentation of a precursor prior to the masker-signal stimulus can be used to infer the time- and level-dependence of the efferent effect.

Previous psychophysical studies examining the decay characteristics of the efferent effect have varied either precursor level or post-precursor delay, but not both together. The present study varies both variables to investigate each separately and also their interaction. The use of the FDMC technique ensures that the effect of the precursor on gain can be measured independently of any masking effects by the precursor. This has been a confound in previous behavioural studies.

**Methods**

FDMCs were obtained from five listeners with and without a precursor noise (500-ms, 1-kHz wide noise band centered at 4 kHz). The precursor was presented at levels of 20, 40, 60 and 80 dB SPL, with silent intervals between precursor and masker (P-M interval) of 0, 50, 100 and 200 ms. A 2-way ANOVA showed a significant effect for precursor level \([F(3,12) = 14.09, p < 0.001]\), P-M interval \([F(3,12) = 5.28, p = 0.015]\) and interaction between precursor level and P-M interval \([F(9,36) = 2.94, p = 0.01]\).

**Results**

Estimated gain decreased as precursor level increased, and for a given precursor level, gain increased as the P-M interval increased, consistent with a decay of the efferent effect.

**Conclusion**

It appears that effect of P-M interval on recovery of gain increases with precursor level.
The Effect of Preceding Tones on Frequency Discrimination
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Background
Demany and colleagues have suggested that the auditory system contains automatic frequency-shift detectors (FSDs) that bind successive sounds together. These hypothetical units appear to be tuned to optimally detect frequency shifts of around 1.5 semitones. The present work details six experiments that may provide more evidence for the existence of FSDs.

Methods
All the experiments used the same basic procedure. Each trial contained a sequence of three pure tones, referred to as (from earliest to latest) the ‘precursor’, the ‘standard’, and the ‘test’ tones. Listeners discriminated frequency shifts between the standard and the test tones ($\Delta_{2}$); the directions of these shifts were randomly upward or downward, and their magnitudes were fixed to the same point on each listener’s psychometric function. Listeners were told to always ignore the frequency shifts between the precursor and the standard tones ($\Delta_{1}$), whose values ranged from 0 to 9 semitones, and whose direction on each trial was independent of the direction of $\Delta_{2}$. Sensitivity ($d'$) and bias ($c$) were measured as a function of $\Delta_{1}$.

Results
In almost all conditions and experiments, there was a consistent nonmonotonic relationship between $d'$ and $\Delta_{1}$. Sensitivity declined as a function of increasing $\Delta_{1}$ up to approximately 1.5 semitones, and further increases in $\Delta_{1}$ increased sensitivity. The ‘dip’ in $d'$ when $\Delta_{1}$ equaled 1.5 semitones was observed even though the precursor and the standard tones were separated by a 500-ms interval, meaning that the results are unlikely to be caused by masking. The dip was observed when listeners judged the direction of frequency shifts and when they simply detected frequency shifts. Neither altering the timbre nor the perceived spatial location of the precursor tones so that they sounded perceptually dissimilar to the remaining tones removed the dip. The dip was not influenced by whether the frequency of the standard tone was roved or fixed over trials, nor by whether the value of $\Delta_{1}$ was random or blocked for 100 sequential trials. The relationship between $c$ and $\Delta_{1}$ was less consistent across listeners and across experiments.

Conclusion
FSDs provide a convenient explanation for the present results if one assumes that (a) listeners discriminate small frequency shifts using FSDs, and (b) FSDs were activated—and consequently fatigued by—the irrelevant frequency shift on each trial. Thus, maximal fatigue would have occurred on trials in which $\Delta_{1}$ equaled 1.5 semitones. [Work supported by NIH R01 DC05216]

Neural Correlates of Auditory Streaming in the Human Brainstem
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Background
Neural correlates of auditory streaming have been reported both in subcortical and cortical sites. However, convincing evidence for neural correlates of streaming in the human brainstem is still limited. Here, we examined the correlation between the auditory brainstem frequency-following response (FFR) and behavioral reports of perceived streaming for a prolonged exposure to a repeated acoustic pattern that may evoke perceptual bistability between a single coherent stream and two segregated streams.

Methods
The stimulus was a repeated triplet-tone sequence (ABA-ABA-....; A: 315-Hz tone, B: 400-Hz tone, -: silence). The duration of each tone was 50 ms, which included rising and falling cosine ramps of 10 ms. The duration of silence was 60 ms. The frequency difference between A and B tones were four semitones. Fifteen normal-hearing adults aged 19 - 34 participated in the experiment. The experiment consisted of 48 sessions. In each session, participants were presented with 200 repetitions of ABA- triplets. While listening, they pressed a button corresponding to one-stream or two-stream percepts whenever they experienced perceptual switching. At the same time, the FFR was recorded using a vertical one-channel electrode montage. The electrode placements were Cz (active), ipsilateral earlobe (reference), and forehead (ground). The recorded FFR was segmented at every onset of ABA- triplets and classified according to behavioral reports (one-stream or two-stream), then averaged within each perceptual category. The results were evaluated in terms of the
amplitude of the average FFR at the stimulus frequencies. The phase locking value (PLV) was also examined. To derive the PLV of a given tone, first, the difference between fourie-transformed stimuli and corresponding FFR was computed for each tone presentation and expressed as an unit vector on the complex plane. Then, the PLV was determined as the length of the vector average across presentations.

Results
The averaged FFR amplitude and PLV to the second A tone (A2) in the triplet were significantly smaller for one-stream than two-stream percepts. For other tones, no significant change was observed between one-stream and two-stream percepts. Moreover, when participants perceived one stream, the averaged FFR amplitude and PLV to A2 were significantly smaller than those to A1. When participants perceived two streams, these differences were not significant.

Conclusion
The results demonstrated significant changes in the FFR according to perceived streaming for the physically constant stimulation, indicating neural correlates of auditory streaming in the human brainstem.

Effect of Phase Curvature on Forward and Backward Masking: Evidence Against Efferent Activation
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Background
Carlyon and Datta (1997) measured forward masking of a 1-kHz signal by Schroeder-phase harmonic complexes. The positive Schroeder-phase masker caused the auditory filter centred on 1-kHz to have a highly modulated output and produced less masking than a negative Schroeder-phase masker, which produced an output with a flatter envelope. This was attributed to fast-acting compression in the auditory system attenuating the peaks of the more modulated auditory filter output. Wojtzack and Oxenham (2009) obtained similar results for a 1-kHz signal frequency (fs) with masker components between 400-1600 Hz, but no phase effect when the masker components were all below fs. They attributed this to the portion of the basilar membrane tuned to 1-kHz responding linearly to the off-frequency masker. However, for fs=6-kHz, a masker phase effect was observed for masker components below fs. They suggested that this off-frequency masker may cause the efferent system to attenuate the response to the 6-kHz probe.

Methods
Experiment 1 replicated Wojtzack and Oxenham main findings. For fs=1-kHz the forward masker contained consecutive equal-amplitude harmonics of 100 Hz: 400–1600 Hz (on-frequency) or 100–600 Hz (off-frequency). The phase of the nth harmonic equalled Cπn(n-1)/N where C=-1 or 1. For fs=6-kHz the on- and off-frequency maskers spanned 4800–7200 Hz and 1600–4000 Hz, respectively. C was -1 or 0. Experiment 2 measured backward masking for fs=6-kHz with the off-frequency masker.

Results
For Experiment 1, on-frequency maskers showed phase effects at both fs, while off-frequency maskers showed a phase effect only with fs=6kHz. For Experiment 2 a substantial phase effect was observed for most listeners, and, across all six listeners tested, the size of this effect correlated highly significantly with that in the analogous forward masking condition (r=0.97, p<0.001). A control forward-masking experiment with a long masker-signal gap showed no masking, ruling out the possibility that, in experiment 2, the backward masker in one interval affected detection of the signal in the previous interval.

Conclusion
The results of both experiments are consistent with fast-acting compression in the auditory system reducing the ‘effective level’ of stimuli that produce ‘peaky’ auditory filter outputs. The 1-kHz data are consistent with compression rising entirely in outer-hair-cells, while the phase effect observed for fs=6-kHz and off-frequency maskers suggest an additional effect (possibly compression in inner-hair-cells, auditory nerve fibres or brainstem). Experiment 2 showed that phase effects with off-frequency maskers are not caused by efferent activation, which cannot act retrospectively.
The effect of cochlear nonlinearities on binaural masking level differences

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Background

The binaural masking level difference (BMLD) has been shown to be constant (10\textendash15\text{dB}) for masker spectrum levels from 70\text{dB}/Hz down to 30\textendash40\text{dB}/Hz and to gradually decrease with lower levels (McFadden, 1968; Hall and Harvey, 1984). The decrease at low levels was larger in an asymmetric condition where the masker was attenuated in only one ear. McFadden predicted the data by assuming that an external and an internal noise would interaurally decorrelate the internal representations of the stimuli. In the present study, the role of nonlinear cochlear processing and asymmetric masker level on the BMLD was investigated using an equalization-cancelation (EC) based binaural model framework.

Methods

The BMLD was measured for 500\text{Hz} target tones presented in 3\text{kHz}\textendashwide maskers. BMLDs were obtained as a function of masker level in one symmetric and two asymmetric masker conditions: (i) \text{NoS\pi}: thresholds were measured for masker spectrum levels between 50 and -10\text{dB}/Hz, with the same level at both ears; (ii) \text{No'S\pi'50}: same as first condition but the masker was attenuated in one ear only and fixed at 50\text{dB}/Hz in the non\textendashattenuated ear; (iii) \text{No'SI'I'20}: same as second condition but with a masker level of 20\text{dB}/Hz in the non\textendashattenuated ear. An EC based binaural model with a frontend including nonlinear peripheral processing (Jepsen et al., 2011) was used to predict these results.

Results

The BMLD obtained in the \text{No'S\pi'50} condition was smaller than that obtained in the \text{NoS\pi} condition at all masker levels between 50 and -10\text{dB}/Hz. The difference in BMLD between the two conditions gradually increased with decreasing masker level from 50 to 20\text{dB}/Hz and remained constant between 20 and -10\text{dB}/Hz. The proposed model could account for these data. A model analysis suggested that the increase in BMLD difference between the \text{No'S\pi'50} and \text{NoS\pi} conditions results from a decrease in interaural correlation at the output of the periphery, and is a consequence of the cochlear nonlinearity at levels between 20 and 50\text{dB}/Hz. For levels below 20\text{dB}/Hz, cochlear processing becomes linear and the model predicts that the difference in BMLD between the symmetric and the two asymmetric conditions is constant, in line with the experimental data.

Conclusion

A model was proposed to account for the effect of level asymmetry on BMLD. The modeling results suggest that cochlear nonlinearities affect the analysis of binaural cues at higher processing stages such that signals carrying interaural level differences can become interaurally decorrelated.
The Audiogram Fails to Reveal Deficits from Significant Inner Hair Cell Loss in Chinchillas Treated with Carboplatin
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Background
The audiogram is the most widely used clinical measure of hearing. It has been useful in detecting and describing acquired hearing loss as a consequence of ageing, a history of noise exposure, otologic disease and/or exposure to ototoxic drugs that preferentially damage the outer hair cells of the inner ear. These hearing losses lead to increased thresholds and poorer tuning. In contrast, despite their extensive innervation of afferent auditory nerve fibers (>90%), little is known about how loss of inner hair cells impacts auditory perception and it is not clear if the audiogram is useful in detecting sensory deficits associated with selective inner hair cell loss. Previous physiological studies in chinchillas with carboplatin induced selective inner hair cell loss suggest reduced compound action potentials at the level of the cochlea but near normal potentials at the inferior colliculus and auditory cortex. The purpose of the present study was to determine if the audiogram could reveal deficits associated with carboplatin induced inner hair cell loss in the chinchilla.

Methods
To determine if selective inner hair cell loss impacts the audiogram, chinchillas were first trained to respond to tonebursts across a broad frequency range (250-11,300 Hz) using an established avoidance paradigm. Subjects were then treated with a single 75 mg/kg dose of carboplatin, a dose known to induce 30-90% inner hair cell loss and reevaluated. Finally subjects were sacrificed and the extent and pattern of inner hair cell loss was correlated with post-carboplatin audiometric data.

Results
Chinchillas treated with carboplatin showed selective inner hair cell loss of 30-90%; results consistent with previous reports. Relatively large lesions (>80%) were needed before a significant threshold shift could be observed. However, this effect was only evident at the higher frequencies with lower frequencies showing no changes and was present in about half the animals.

Conclusion
These data suggest that audiometric threshold assessment does not reveal the presence of inner hair cell lesions even when these extend over a large region of the cochlea. We suggest that conventional audiology is likely insensitive to inner hair cell loss in humans as well. Only small populations of inner hair cells appear to be necessary for detecting tone stimuli in a quiet background.
Auditory Filter Bandwidths in the Common Marmoset (Callithrixjacchus)
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Background
The common marmoset has emerged as a promising non-human primate model in auditory neuroscience. However, a full appreciation of the physiological response to sound will require a basic understanding of auditory perception and behavioral sensitivity in this species, including measures of auditory frequency selectivity.

Methods
We measured frequency selectivity in marmosets by estimating auditory filter bandwidths using a psychophysical task in which tone thresholds were measured as a function of notched noise masker bandwidth. The resultant threshold data was used to estimate the auditory filter equivalent rectangular bandwidths (ERBs) using procedures originally developed for human subjects. We tested five different frequencies spanning much of the hearing range of this species (250 Hz, 500 Hz, 1000 Hz, 7000 Hz, and 16000 Hz).

Results
Marmosets have ERBs that are generally comparable to those measured in other mammalian species, although ERBs in marmosets tend to be wider than those measured in humans. This was true for all frequencies tested except 7000 Hz, where marmoset filter bandwidths are narrower than those measured in humans. Wider bandwidths likely means that, at most of the frequencies tested here, marmosets have poorer spectral resolution compared to humans. Narrower filter bandwidths at 7000 Hz probably reflects the fact that this is approximately the frequency of best hearing in this species and also the frequency around which most of the spectral energy in marmoset vocalizations is concentrated.

Conclusion
Differences in the size of human and marmoset filter bandwidths may reflect a series of tradeoffs between basilar membrane length (which is half that measured in humans), hearing range (which extends about 10 kHz higher than in humans), and the need to process species-specific communication signals centered in a particular range of frequencies. [Research supported by NIH grants DC003180, DC005808]

Hcn1 null mutant mice show delays in processing sounds on time scales from 1 ms to over 2 seconds in tasks that provide animal models for speech encoding, temporal summation, and forward masking.
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Background
In vitro studies of auditory brainstem neurons demonstrate that their precise timing depends on a low input resistance that gives them a short time constant (Oertel, 1983), and that this feature depends in part on a hyperpolarization-activated, mixed cation conductance (gh) in channels that contain Hcn1 subunits (Cao & Oertel, 2011; Kopp-Scheinpflug et al. 2011).

Methods
Here we examine the functional significance of this conductance in behavioral comparisons of Hcn1 +/- mice (n = 63) and Hcn1 -/- mice (n = 60) using standard startle reflex (ASR) modification measures to determine: (1) their threshold for gap detection; (2) temporal summation of near-response-threshold tone pips; and (3) the rate of recovery from (a) suprathreshold gaps and (b) from suprathreshold noise bursts in a refractory paradigm. We measured also their ABR thresholds and suprathreshold amplitudes.

Results
Gap detection thresholds were delayed by 1 ms in -/- mice, but they had higher and more persistent response suppression from suprathreshold gaps than +/- mice: the between group difference began at a 30 ms interval after the gap and lasted for over 100 ms. The peak of 2-pulse temporal summation was delayed in -/- mice from 2 to 3 ms, with +/- mice responding more at short delays but less at longer delays of 3 to 4 ms. The refractory (suppressive) effect of one startle stimulus on the response to the second stimulus was more persistent in -/- mice, with the groups being different at 1 and 2 second recovery times, but not at 4 seconds. ABR thresholds in young adults (median age ~ 60 PND) were equal up to 24 kHz, but -/- mice showed a hearing loss of ~20 dB at 32 and 48 kHz. P1 for clicks had near identical latencies (means of 1.46 and 1.43 ms), but the -/- mice had smaller P1 amplitudes (and also a smaller baseline ASR). The -/- mice
also had a significantly smaller peak amplitude about 1 ms after P1 but higher levels of asynchronous activity beyond 4 ms.

Conclusion
These in vivo behavioral and electrophysiological findings extend the in vitro data in showing that the Hcn1 gene contributes to temporal processing for complex auditory stimuli, but in addition, that it must contribute also to very high frequency hearing.

Physiologically-based Envelope Cues for Diotic and Dichotic Tone-in-noise Detection
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Background
The goal of studying masked detection is to understand what cues listeners use to detect signals in noise. Envelope cues derived from the stimulus predict detection performance successfully for both diotic and dichotic conditions. Here, it was hypothesized that predictions of detection performance based on envelope cues derived from physiological models would be similar to predictions using stimulus-based envelope cues.

Methods
500-Hz tones of 300-ms duration were detected in 300-ms narrowband and wideband reproducible Gaussian noises. Model predictions of the variance in detection performance across a set of reproducible noise waveforms were computed for both diotic and dichotic conditions.

The stimulus-based envelope cue for the diotic condition was computed from a modified envelope-slope (ES; Richards, 1992, JASA 91:3424; Davidson, et al., 2006 JASA 119:2258) model. A bandpass filter extracted the 40 to 140 Hz envelope frequency range, which contained the most information about tone presence. For the dichotic condition, the slope of the interaural envelope difference (SIED) model, which examined the difference between the envelopes at the two ears, was used to predict listeners' detection performance.

For the physiologically-based envelope models, the stimulus was passed through a human version of the auditory-nerve (AN) model (Zilany et al., 2009, JASA 126:2390; Ibrahim and Bruce, 2010, The Neuropsychological Bases of Auditory Perception, pp. 429) to obtain the synaptic response. Model AN responses were used as inputs to a same-frequency inhibition and excitation model for the cochlear nucleus (Nelson and Carney, 2004, JASA 116:2173). A bandpass modulation filter simulated an amplitude-modulation tuned inferior colliculus (IC) cell. Envelope-slope information was calculated as the derivative of the modulation filter response. The model decision variable was the integral of envelope slope over time.

Results
For the diotic condition, predictions based on the model IC response were similar to those of the modified ES model. For the dichotic condition, predictions based on binaural physiological envelope cues were similar to those of the SIED model. Moreover, for the dichotic wideband waveforms, predictions from both the stimulus-based and physiological envelope models were better than those of other dichotic models.

Conclusion
Physiological models for processing envelope cues predicted a similar amount of the variance in both diotic and dichotic detection performance as the stimulus-based envelope cues. Thus, known physiological mechanisms are feasible for extracting and using monaural and binaural envelope cues for detection.

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Discrimination of Partial USVs by CBA/CaJ Mice
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Background
Mice are a commonly used model in hearing research, yet little is known about how they perceive their own ultrasonic vocalizations (USVs). Recent studies illustrate that mice are poor at discriminating between some of these vocalizations (Neilans et al. ARO 2013 abstract). For example, discrimination accuracy in mice is low for calls that are reversed in time, which is surprising when compared to similar studies in humans and other animals. This leads to the question of how
much information the mice need from their calls in order to identify them and which portions of the USVs are most important for recognition. To answer these questions, the present study looks at how well mice can discriminate targets containing small portions of a call from a repeating background containing the whole call. Based on human and bird studies, we hypothesized that discrimination performance would be higher as more of the call was deleted, and that there would be an asymmetry in discrimination abilities depending on whether the beginning or the end of the call was omitted.

**Methods**
CBA/CaJ mice were trained and tested in a discrimination task using operant conditioning procedures. The mice were required to discriminate target stimuli from a background call to receive water/chocolate milk reinforcement. The target stimuli were incomplete versions of the repeating background stimulus, containing either one-third (initial third, middle third, and last third) or two-thirds (initial two-thirds, last two-thirds, and middle third removed) of the whole call. The first twenty responses for each target stimulus were recorded and analyzed.

**Results**
The discrimination results show that performance was very low for all targets. Discrimination accuracy did not improve with decreasing durations (two-thirds versus one-third), and results were similar across every partial-USV target. The mice could not discriminate small portions of the calls from the whole call.

**Conclusion**
The current results suggest that different portions of the call do not contain any more information than other portions of the call, contrasting with past results from humans and birds. This indicates that mice maybe perceiving their vocalizations differently from humans and birds. However, these calls are in the ultrasonic range and are shorter in duration than any human or bird vocalizations and thus may be difficult to compare.
Enhancement and Suppression Estimated from Growth of Masking Functions
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Background
Threshold for a signal in a simultaneous masker may decrease when the signal is delayed from masker onset or when a precursor is added. When the masker has energy at the signal frequency, the results have been modeled as a decrease in cochlear gain, which could be consistent with the medial olivocochlear reflex (MOCR). When the masker lacks energy at the signal frequency, the results are consistent with a decrease in suppression, but this has not been directly demonstrated. One way to estimate suppression is to compare thresholds for a signal in simultaneous and forward masking using a masker that is approximately an octave below the signal frequency. Because the masker response at the signal place should be linear, the difference in thresholds between the two conditions yields the amount of suppression of the signal in dB. In the present experiment, enhancement and suppression were measured from the same data.

Methods
Ten subjects with normal hearing were tested. The signal was 4 kHz and 6 ms in duration. Maskers were narrowband noise centered at 2.4 kHz (off-frequency) and 4 kHz (on-frequency). Growth of masking functions were measured with masker level fixed and signal level varied. In the baseline condition, the masker duration was 20 ms. The signal immediately followed the masker or was presented in the center of the masker. For the comparison condition, the duration of the simultaneous masker was lengthened by 40 ms prior to signal onset, or a separate, fixed-level precursor was presented immediately before the masker.

Results
Enhancement was calculated for a fixed masker level as the difference in signal threshold for the short simultaneous masker and the long simultaneous masker or the precursor + masker. To calculate suppression, the masker level necessary to mask a fixed signal level was estimated from the data. Suppression was calculated as the masker level in the forward masking condition minus threshold in the simultaneous masking conditions. Suppression was greatest for the short masker and decreased in the long masker or the precursor + masker conditions. The amount of enhancement depended on the amount of decrease in suppression and the slope of the input/output function in the region of the signal level.

Conclusion
Results suggest that for these conditions, enhancement is consistent with a decrease in suppression coupled with the effects of compression.

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Survival and integration of mouse inner ear progenitor/stem cells in the cochleas of hearing-impaired guinea pigs

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Background
In mammals, damage to the sensory receptor cells (hair cells) of the inner ear results in permanent sensorineural hearing loss. Here, we investigate whether postnatal mouse inner ear progenitor/stem cells (mIESCs) can survive and integrate after transplantation into the basal turns of neomycin-injured guinea pig cochleas. We also studied the potential effects of the cell transplantation on auditory function.

Methods
Eight adult guinea pigs were deafened by intratympanic neomycin delivery. After 7 days, the animals were randomly divided in two groups. The study group (n=4) received a transplantation of 1 X 10⁵ LacZ-positive mIESCs in culture media to the scala tympani. The control group (n=4) received culture media only. Fourteen days after the transplantation, functional analyses were performed by auditory brainstem response (ABR) measurement, and the animals were sacrificed. The presence and distribution of mIESCs were evaluated by immunohistochemistry of longitudinal sections of the cochlea from the study group. Non-parametric tests were used for statistical analysis of the data.

Results
Intratympanic neomycin delivery damaged hair cells and increased auditory thresholds prior to cell transplantation. There were no significant differences between auditory brainstem thresholds before and after transplantation in individual guinea pigs. Some mIESCs were observed in all scalae of the basal turns of the injured cochleas, and a proportion of those cells expressed the hair cell marker myosin VIIa. Some transplanted mIESCs integrated in the cochlear basilar membrane. There was no evidence of inflammatory infiltration in any of the guinea pigs’ cochleas.

Conclusion
Although mIESC implantation resulted in no obvious effect on auditory thresholds, our experiments demonstrated the survival, migration, integration and expression of a hair cell marker in transplanted cells.
Treatment for Connexin30 Deletion Associated Hearing Loss Utilizing Otocyst Trans Uterine Gene Transfer in Mice

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Background
Mutation in gap junction beta-6 (GJB6), the gene that codes for Connexin30 (Cx30), causes hereditary deafness in humans and mice. In our previous study, we show that electroporation-mediated transfection of Cx30-targeted shRNA (shRNA-Cx30) into the normal mouse otocyst induced the knockdown of endogenous Cx30 in the cochlea, lack of endocochlear potential (EP) and severe hearing impairment. Now then, we investigated whether trans-uterine gene transfer therapy could cure hearing loss in Cx30 knockout mice.

Methods
At embryonic day 11.5 (E11.5), EGFP-fused Cx30 plasmid vector was microinjected through the uterus into the otic vesicle of Cx30 knockout mice and electroporated. Electroporated embryos were delivered at E18.5. Some fetuses were removed to prepare frozen sections, and some fetuses were raised by surrogate mothers until functional and morphological assessments on postnatal day 30 (P30). An auditory brainstem response (ABR), endocochlear potential (EP), immunohistology were used for the assessments. As a control, EGFP plasmid was used.

Results
Immunohistological analyses revealed that Cx30-positive cells were detected in the medial walls, the prosensory lesions, the lateral walls and the spiral ganglions at E18.5; they were detected in the spiral limbuses, organs of Corti, stria vascularises, spiral ligaments and spiral ganglions in the treated ears at P30. There was no Cx30 expression in the non-treated right ears. The treated side ears exhibited significant amelioration of the auditory threshold compared with the non-treated right ears. (n=4, p<0.02). The treated ears showed normal EPs (n=2).

Conclusion
Cx30-deficient homozygous mice exhibit severely impaired hearing and lack endocochlear potentials. Trans-uterine transfer of normal Cx30 genes into the otocysts of Cx30-deficient homozygous mice reversed the downregulation of Cx30 in the cochleae, and thereby restored their auditory functioning. Trans-uterine gene transfer into the otocyst is a novel treatment for genetic hearing loss.
Combined Expression of MicroRNAs and Transcription Factors for Promoting Hair Cell Differentiation

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Background
Damage to mechanosensory hair cells (HCs) of the inner ear causes permanent hearing loss in millions of people each year. Although some vertebrate species can regenerate HCs, human auditory HCs lack regenerative capacity. An emerging paradigm in regenerative strategies is to utilize a combination of crucial factors to achieve cell reprogramming. The transcription factor (TF) Atoh1 is indisputably required for HC development and represents the standard for regenerative strategies. However, microRNAs (miRNAs) post-transcriptionally regulate gene expression and are also required for HC differentiation and maintenance. These include miRNA-183 family members miR-183, miR-96, and miR-182. Additionally, other TFs including Pou4f3 and Gfi1 are necessary for hair cell differentiation and survival. Our objective is to apply a combination of these crucial factors to hair cell generation.

Methods
To determine whether a combination of miRNAs and TFs can generate HCs more effectively than Atoh1 alone, we have developed a novel vector for expression of miRNAs and multiple TFs from a single transcript. The transcript produces miR-183 family members from an intron, and individual TFs (Atoh1, Pou4f3, and Gfi1) and red fluorescent protein (RFP) from a single open reading frame by ribosomal cleavage of intervening viral peptide elements. This parent construct was used to generate a series that contain different combinations of factors, and certain constructs were converted to adenovirus vectors (AdVs) for gene delivery. miRNA and protein expression was examined in mouse otic precursor cells (IMO-2B1) by quantitative PCR (qPCR) and fluorescence microscopy. AdVs were used to infect mouse induced pluripotent stem cells (iPSCs), and development of HC-like morphology was assessed by fluorescence microscopy.

Results
Elevated expression of miR-183 family members in IMO-2B1 cells was observed by two different qPCR methods for mature miRNA detection. RFP expression in IMO-2B1 cells observed by fluorescence microscopy is indicative of factor expression given that RFP is produced from the final open reading frame of each construct. However, vectors encoding more factors yielded less RFP expression than those with fewer factors. Preliminary assessment of AdV effects on iPSCs using f-actin staining showed that the combination of miR-183 family and Atoh1 produced morphologically HC-like cells. In comparison, Atoh1 alone marginally affected iPSC cytomorphology.

Conclusion
Preliminary results suggest that combining HC-specific miRNAs and TFs provides a more effective means for cell reprogramming and HC differentiation that might be useful in therapeutic strategies for HC repair and regeneration.
Development of Acoustic Sensor for Wide-Range Frequency Selectivity Using Non-Uniform Thick Ladder-Like BAM

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Background

The basilar membrane in a cochlea has an important role to realize the frequency selectivity. In previous study \cite{1}, the authors developed a bionic auditory membrane (BAM) which mimics cochlear functions and proposed to use it as a sound processor. Although the basic applicability of the BAM was confirmed, the frequency range in which the BAM could work was narrower than the human audible range. One of the reasons is the uniform thickness of the BAM, though the thickness of the basilar membrane is varied by location. Therefore, we have newly developed a non-uniform thick ladder-like BAM. While the previous BAM has a trapezoidal membrane structure, the device is designed as an array of beams.

Methods

The non-uniform thick BAM was fabricated by the grayscale lithography based on MEMS (microelectromechanical systems) processes. The vibration characteristics of the BAM in air was measured as a preliminary experiment and evaluated the efficiency of the thickness change. The vibrations of beams were measured by laser Doppler vibrometer. Since the cochlea is filled with lymph fluids, the vibration characteristics of the BAM immersed in liquids were also measured. Here, the BAM was fixed in a liquid filled channel and the acoustic waves were applied by a piezoelectric actuator.

Results

The BAM consists of 64 micro beams and the thicknesses are successfully changed from 3.02 \(\mu\)m to 139 \(\mu\)m. The length and the width of beams are also changed from 750 \(\mu\)m to 1500 \(\mu\)m and from 50 \(\mu\)m to 600 \(\mu\)m, respectively. The frequency range is quantitatively evaluated using the ratio \(f_{\text{max}}/f_{\text{min}}\) between the maximum first mode resonant frequency \(f_{\text{max}}\) and the minimum one \(f_{\text{min}}\). The range in the non-uniform thick BAM is 10.9 times wider than that in the uniform one, indicating the successful control of the frequency range by the thickness change. The result in liquid shows that the resonant frequency is decreased by the fluid-structure interaction compared with that in air. From this effect, the ratio in liquid is 3.13 times larger than that in air.

Conclusion

Although the resonant frequency of the BAM is still higher than the human audibility, these experimental results prove the feasibility of the new BAM as an acoustic sensor which can work over the full range of the human audible frequency.

Development of a Selective Hair Cell-Spiral Ganglion Neuron Double Ablation Model to Study Cochlear Implantation in Combination with Stem Cells

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Background
Animal models that resemble lesions observed in clinical conditions are of great value to explore future therapies. We are interested in developing the potential applications of stem cells for the treatment of hearing loss, particularly in combination with cochlear implants. We have previously employed the gerbil auditory neuropathy model developed by Lang et al. (JARO 6; 63-74, 2005) to study the ability of hESC-derived otic neuroprogenitors (hONPs) to repopulate the damaged spiral ganglion and to restore auditory function. After topical application of ouabain to the round window, permanent auditory neuron degeneration is induced, leading to a rise of ABR thresholds in response to click and tone stimuli. After an infusion of hONPs into the modiolus, we have evidence that the transplanted cells can graft and differentiate into bipolar, βIII-tubulin positive neurons which contact the hair cells and project to the brainstem. Moreover, we have obtained a significant improvement in the mean auditory thresholds. This model shows preservation of the DPOAE responses, demonstrating that the hair cells remain intact.

Methods
We are currently exploring a 'double ablation' model, in which aminoglycoside antibiotics are used to abrogate hair cells alongside the ouabain-induced neuropathy. We have explored several paradigms of aminoglycoside application, with variable degrees of success.

Results
On one hand, application of gentamicin or kanamycin directly to the round window, or systemic kanamycin followed by the diuretic bumetanide does not induce hair cell loss. Alternatively, the combined application of kanamycin and furosemide is very effective in killing the hair cells, substantially raising ABR thresholds. In this condition however, the spiral ganglion neurons are preserved even months after aminoglycoside application, resembling the variable preservation of neurons after hair cell damage observed in humans.

Conclusion
The ability to damage selectively hair cells or neurons adds flexibility to a model system for studying stem cell transplantation. Moreover, the combined application of aminoglycosides and ouabain should prove ideal to study the effects of a fully-implantable, cochlear stimulator to substitute for hair cell function combined with the stem cell-replacement of spiral ganglion neurons, thus facilitating the application of cochlear implants to a broader range of patients.
Partial Regeneration Following Cisplatin Ototoxicity in Zebrafish Lateral Line

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Background
Cisplatin is a chemotherapeutic agent that is used to treat a variety of solid tumors. Ototoxicity is a frequent side effect, often resulting in permanent sensorineural hearing loss. Nonmammalian vertebrates are capable of regenerating hair cells after aminoglycoside injury, but the avian ear cannot regenerate after cisplatin ototoxicity (EL Slattery, ME Warchol, J Neurosci, 2010). The current study analyzed hair cell death and regeneration in the zebrafish lateral line after cisplatin administration.

Methods
Zebrafish larvae (5 days post-fertilization) were treated for 4 hours in 1000 µM cisplatin and then allowed to recover for 1-10 days. Other fish were exposed to cisplatin, followed by treatment for 24-72 hours with the γ-secretase inhibitor DAPT (50 µM). After fixation, specimens were immunolabeled for HCS-1, in order to identify hair cells. Hair cells were quantified from 10 individual neuromasts along the posterior lateral line of each fish.

Results
We first examined hair cell numbers within three specified neuromasts evenly distributed along the length of the posterior lateral line. We found that neuromasts of unexposed (control) fish contained 9.3±3.4 HC/neuromast. However, at 1 day after cisplatin exposure, hair cell numbers had decreased, to 1.7±0.5 HC/neuromast. Quantification of hair cells numbers at 10 days after cisplatin revealed that neuromasts now contained 5.5 ±3.0 HC’s (p<0.0001). Treatment with DAPT for 72 hours after cisplatin exposure resulted in only a very slight increase in hair cell numbers, compared to controls.

Conclusion
Consistent with previous studies (e.g, H Ou et al., Hearing Res, 2007), our data indicate that treatment for 4 hours in 1000 µM cisplatin results in the death of most hair cells within neuromasts of the posterior lateral line. Additionally, our data show that the zebrafish lateral line can regenerate hair cells after cisplatin ototoxicity. Previous studies of regeneration after aminoglycoside ototoxicity (e.g., JA Harris et al., JARO, 2003; EY Ma et al., J Neurosci, 2007) indicate that hair cell recovery is complete within three days of injury. However, our data show that regeneration after cisplatin ototoxicity is still incomplete after 10 days recovery. In addition, inhibition of Notch signaling (via treatment with DAPT) does not result in a substantial increase in hair cell numbers. Since hair cell regeneration is mediated by epithelial supporting cells, our observations suggest that cisplatin may have toxic effects on both hair cells and supporting cells.
Hearing improvement by gene therapy in a mouse model created by a conditional knockout of Gjb2 gene

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Background
Hereditary deafness affects about 1 in 2,000 children and mutations in the GJB2 gene, coding the gap junction, are the major causes in various ethnic groups, which require normal gene transfer in the early developmental stage to prevent deafness. Mice present an ideal model for inner ear gene therapy because their genome is being rapidly sequenced and their generation time is short. In order to establish the fundamental therapy of congenital deafness, we generated targeted disruption of mouse Gjb2 gene using Cre recombinase controlled by P0. Using this animal model, we examined the potential of gene therapy in the inner ear, using the homozygous mutant mice and the heterozygous mutant mice.

Methods
Adeno-associated virus vectors (AAV) carrying Gjb2 gene were injected into the scala tympani through the round window of the cochlea of the homozygous mutant adult and neonatal mice.

Results
In the adult mice, the expression of Cx26 was observed in the fibrocytes of the spiral ligament and spiral limbus, but was not seen in the supporting cells and failed to improve the hearing ability. However, in the neonatal mice, the expression of Cx26 was seen in the supporting cells and the hearing ability was improved. In the histological analysis, the organs of Corti of the homozygous mutant mice injected AAV carrying Gjb2 gene were higher than the non-injected homozygous mutant mice.

Conclusion
The present paper will present the data regarding introduction of the virus vector into the Gjb2 knockout mouse at the neonatal stage.
Cochlear gene transfer mediated by adeno-associated virus: comparison of two surgical approaches

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Background
Hearing loss is one of the most common disabilities affecting the US population. Gene therapy offers the possibility of delivering corrective genes to the cochlea, potentially improving hearing. Currently, the most commonly used surgical methods for viral gene therapy delivery to the cochlea are the round window and the cochleostomy approaches. However, the patterns of viral infection and the effects on hearing have not been directly compared between these two approaches. In this study, we compare the patterns of cochlear infection and effects on hearing between these two surgical approaches using adeno-associated virus serotype 8 (AAV8) as the gene delivery vehicle.

Methods
CBA/J mice were used in this study. AAV8-GFP was delivered to the cochlea by either the round window or the cochleostomy approach. Auditory brainstem-evoked response (ABR) was used to examine hearing thresholds before and after surgery. Animals were examined at 1, 2, 3, and 4 weeks after surgery for the patterns of cochlear infection and hearing loss.

Results
Cochlear gene transfer was successful through both surgical approaches. In both approaches, the inner and outer hair cells, supporting cells, and the striavascularis were infected with AAV8-GFP. The cochleostomy approach caused a greater amount of hearing loss than the round-window approach.

Conclusion
The round window approach and the cochleostomy approach are both successful at AAV-mediated gene transfer to the cochlea. The round window approach resulted in less hearing loss compared to the cochleostomy approach.
Essential role for CBP in Tlx3-mediated neuronal differentiation from mouse embryonic stem cells
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Background
The histone acetyltransferase CBP is a known transcriptional co-activator and has been shown to bind homeobox proteins, including Tlx3. The association of CBP with the homeobox proteins appears to be critical for evoking transcriptional activation of their target genes. Using Tlx3-expressing mouse embryonic stem cells (ESCs), we have shown that: (1) Tlx3 regulates Ngn1 expression only after neural induction; (2) Tlx3 binds CBP only after neural induction. We hypothesized that, upon neural induction, CBP may be recruited to and bind Tlx3 occupying the Ngn1 locus, which causes conformational changes in chromatin through acetylation of histone H3/H4 and subsequent transactivation of Ngn1.

Methods
To test which region in Tlx3 is required for CBP binding, we generated several Tlx3 mutants, including Tlx3C lacking the C-terminus domain, and Tlx3ΔHD lacking the homeodomain. We also generated ESC lines stably expressing wild-type Tlx3 and Tlx3ΔHD, which were grown in neural induction medium for 4 or 7 days. To evaluate the functional outcome of impaired Tlx3-CBP binding, we measured changes in intracellular Ca2+ levels.

Results
Immunoprecipitation of HEK293 cells co-expressing CBP and one of these Tlx3 mutants revealed that the homeodomain in Tlx3 is required for its binding to CBP. qRT-PCR and Western blot analyses of samples collected from ESC-derived neurons revealed that Tlx3-CBP binding is essential for proper expression of Ngn1, but not Tau. In addition, a significantly smaller glutamate-induced Ca2+ influx was detected in Tlx3ΔHD-expressing ESC-derived neurons when compared to wild-type Tlx3-expressing neurons.

Conclusion
These results strongly suggest that epigenetic regulation of Tlx3-mediated transcription can modulate efficiency of synaptic transmission in ESC-derived neurons.
Human embryonic stem cell incorporation into the mouse cochlea in vitro
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Background
Hearing loss is one of the most common disabilities affecting the US population. It is often caused by a loss of hair cells and/or spiral ganglion cells in the cochlea. Stem cell therapy offers the possibility of replacing these damaged and missing cells with functional ones and can potentially improve hearing function. One of the major hurdles of cochlear stem cell therapy is the lack of incorporation of transplanted cells into the neurosensory epithelium. In this study, we examine the patterns of human embryonic stem cell (hESC) incorporation into the mouse cochlea in vitro.

Methods
Mouse cochlear explants were dissected from E13, E15, E17, and P0 animals. WA07 hESCs were dissociated into single cells and co-cultured with mouse cochlear explants for 6 days. The patterns of hESC incorporation into the mouse cochlear explants were examined.

Results
Human embryonic stem cells were successfully incorporated into the mouse cochlear explants from E13 and E15 age groups. However, no hESC incorporation was seen when co-cultured with mouse cochlear explants from E17 and P0 age groups. Several hESCs differentiated into neuron-like cells in all cochlear explants examined. Many of these neuron-like cells sent processes to make contact with mouse hair cells. No hair cell-like cells were seen to have differentiated from hESCs in all co-cultures examined.

Conclusion
E13 and E15 mouse cochleae were more susceptible to hESC incorporation into the neurosensory epithelium compared to older age groups. In addition, many hESCs readily differentiated into neuron-like cells and made contact with the mouse hair cells.
Auditory Scene Analysis in Birds: Spectrotemporal Cues Facilitate Streaming
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Background
An animal’s ability to accurately separate sounds from multiple sources in the environment is crucial for its survival. Understanding how different auditory cues in conspecific and heterospecific sounds are perceived by animals and what information facilitates acoustic segregation versus fusion will improve our knowledge of auditory scene analysis. The current study utilizes birdsong to measure streaming in budgerigars (Melopsittacus undulatus) and zebra finches (Taeniopygia guttata). A set of experiments was conducted to 1) determine the role of intensive, spatial, temporal, and spectral cues on streaming; and 2) to establish what specific cue properties are the most important for auditory scene analysis.

Methods
This study uses operant conditioning procedures to determine how conspecific and heterospecific cues affect streaming. Both species were trained to differentially peck keys in response to either a synthetic zebra finch song consisting of five syllables (whole song) or to the same song with one syllable omitted (broken song). Probe trials were presented to the birds on a small portion of all trials where the missing syllable was manipulated. In the first experiment, the original missing syllable was included but (1) varied in spatial location, (2) varied in intensity, (3) occurred earlier or later in time, or (4) varied in its spectral properties from the original whole song. In the second experiment, different stimuli were inserted into the missing syllable location, including a (1) zebra finch call, (2) budgerigar call, (3) budgerigar warble element, and (4) zebra finch syllable not found in the original training song. In both experiments, if the birds respond as hearing a whole song, it implies that the altered content is perceptually fused with the rest of the syllables to form one auditory stream.

Results
Results from Experiment 1 suggest that spatial and spectral cues have the most influence on streaming, while intensive and temporal cues are less informative. In Experiment 2, the probe stimuli that were more likely to be grouped with the surrounding broken song were probes that were the most temporally and spectrally similar to the original syllable and to the surrounding zebra finch song.

Conclusion
These results support the idea that spatial, spectral, and temporal cues are all somewhat important for the streaming of birdsong by zebra finches and budgerigars. These findings are consistent with previous results from humans and other animals.
Aesthetic Perception of Fractal Structure in Melodic Stmuli

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Background

1/f, or fractal, structure is prevalent in nature, biology, and art: landscapes (coastlines, mountains), physiology (heart rate), and visual artwork (Pollock's paintings). Studies have demonstrated that subjects have an aesthetic preference for fractal images when compared to images created with non-fractal structure or random noise. Fractal analysis characterizes long-term correlation and self-similarity by taking sequential aspects of a time series into account.

Methods

A behavioral study was conducted to assess subjects' aesthetic perceptions of melodies with varying degrees of structure. Melodies were created using three types of noise: 1/f (white) contained no long term correlation or fractal structure; 1/f (pink) contained fractal structure and long-term correlation; 1/f (brown) contained highly correlated values. Fourteen adult subjects (musician and non-musician) with normal hearing were instructed to listen to each melody and answer the following questions: 1) Please classify the sequence using a likert scale from 1=ugly to 11=beautiful, 2) Rate the complexity of the sequence (1=least complex to 11=most complex), 3) How musical is the sequence? (1=very un-musical to 11=very musical), and 4) Which type of noise created this melody? Subjects were shown a picture of white, pink, and brown noise.

Results

Results showed that subjects were able to differentiate between white, pink, and brown melodies (p<.01) when asked to rate the complexity of the melody and when asked which type of noise was used to create the melody (p<.01). White melodies were rated as the most complex, pink melodies were the second most complex, and brown melodies were rated as the least complex. Subjects rated white and pink melodies significantly higher than brown on the measures of beauty and musicality (p<.01), showing that subjects preferred white and pink to brown, but did not prefer the pink over white or vice versa.

Conclusion

Music must be predictable enough for people to attend to, but also contain enough violations of expectancy to keep the stimulus novel. The melodies that exhibited white or pink structure, received higher aesthetic ratings than the brown melodies. Possibly because the white and pink melodies were more unpredictable, and therefore more interesting than the brown melodies which were highly repetitive. Researchers have suggested that neural systems have evolved to encode these 1/f signals more efficiently than non-fractal signals; this could explain a conscious, aesthetic preference for 1/f stimuli.
Delayed recovery of auditory stream segregation: cognitive and acoustic disruptions are additive

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Background
People routinely segregate a mixture of sounds into distinct percepts. Previous research indicates that this useful perceptual organization can be disrupted by abrupt changes in auditory scene (Bregman, 1978) or by shifts in attention (Cusack et al., 2004). Here we examined the extent to which switches in attention between tasks and/or abrupt changes in auditory scene (1) cause a segregated percept to "re-set" to an integrated one and (2) influence the amount of time required to recover the segregated percept.

Methods
During each 12s trial, participants were cued to detect targets either in a two-tone (ABA-) auditory sequence for which segregation builds up over time, or in an unrelated distractor task in the opposite ear (Figure 1). Targets in the ABA task were 50ms delays of the B tone 'skips' (Thompson et al, ref) that occurred with 50% likelihood at 3s, 6s, or 9s after trial onset. Target detection sensitivity (d') on the ABA task is much higher when the percept is integrated compared to segregated. The distractor task required listeners to discriminate between two noise types with different amplitude envelopes. Across trials (Figure 2), either the attended task changed (the participant started to attend to the other task in the opposite ear), the auditory scene changed (the stimuli switched ears and the participant 'followed' the stimuli to the contralateral ear), both the task and auditory scene changed simultaneously (the participant kept listening to the same ear, but started doing the other task), or neither changed.

Results
Performance (Figure 3) was analyzed with a 4 (Switch Type) by 3 (Time) MANOVA. Sensitivity early in the trial (3s) indicated that changes in task, auditory scene, or both caused the perceptual organization to "re-set" (high d'). When only the task or auditory scene changed, participants generally showed a return to segregation (low d') by the middle of the trial (6s). However, when both task and auditory scene changed, participants did not show segregation by the end of the trial (9s).

Conclusion
The delayed recovery of segregation following changes in the auditory scene and attended task (even if the attended ear stayed constant) demonstrates an additive effect of cognitive and acoustic disruptions on the buildup of segregation, suggesting that these two disruptions act via different mechanisms.
Recovering Sound Sources from Embedded Repetition And Directed Attention: Effect of Spectral Information on Sound Segregation

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Background

McDermott et al. demonstrated that listeners can identify novel ‘target’ sounds from mixtures if they are presented multiple times across different distractors, even in conditions where single mixtures were impossible to segregate [McDermott, et al. (2011) PNAS. 108(3):1188-93]. Their results indicate that the auditory system can recover sound sources from mixtures by detecting repeating spectro-temporal structure embedded in the acoustic input. In the present series of experiments we aim to investigate whether this repetition-based sound segregation requires selective attention to sounds or whether it can occur partially outside the focus of attention. We adapted the original paradigm of McDermott et al. to a dual task design in which subjects are required to perform a difficult ‘decoy’ task concurrent with presentation of the sequence of sound mixtures.

Methods

Trials consist of ten sound mixtures, each composed of a repeating target and a distractor that changes from mixture to mixture. At the end of a trial, participants judge whether a subsequently presented probe appeared in the sound mixtures (a test of their ability to segregate the sounds in the mixture). Concurrently, subjects are presented with ‘decoy’ visual (Exp1, 2) or auditory (Exp3) stimuli. We manipulate the attentional load of the decoy task by instructing subjects to perform (in different blocks) ‘low load’ and ‘high load’ tasks on these signals. They are instructed (and explicitly motivated) to focus their attention on the ‘decoy’ task, and guess if unsure about the sound-mixture task.

In Experiment 1 we use a rapid sequence of serially presented visual stimuli (digits). In the ‘low load’ condition listeners are required to report whether an underline appeared with one of digits. In the ‘high load’ condition, they are required to memorize the digit order – a task that requires significant attentional and echoic memory resources. In Experiment 2, we use a multiple object tracking visual task, which is attentionally demanding but does not rely on echoic memory. In Experiment 3, we employ a decoy auditory task in which listeners are required to memorize a sequence of rapidly presented sounds.

Results

This study is ongoing. The result of experiment 1 demonstrate that a concurrent, high load, visual task results in similar performance levels to those measured during a low load task.

Conclusion

Our results suggest that listeners can segregate novel sounds from mixtures when attention is directed elsewhere, providing support for repetition-based segregation as automatic, bottom up process.
Perceptual Asymmetry Induced by the Auditory Continuity Illusion
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Background
Listening for messages within mixtures of maskers is both challenging and relevant to daily communication. One aspect of this listening challenge is that masked, or obscured, portions of messages are often perceptually “filled in” to create a continuous stream of information. This auditory continuity illusion has been studied primarily with the stimuli presented in isolation, and it is unknown whether the continuity illusion occurs in more complex situations when attention is not directed towards the interrupted stimulus.

Methods
Young normal-hearing participants listened for target tones in “clouds” of masking tones and noises with pseudo-random spectro-temporal locations. The basic experimental task was to listen for a long (300ms) tone in a cloud of short (100 ms) tones, and its counterpart condition was to listen for a short tone in a cloud of long tones. These two conditions have previously illustrated perceptual asymmetry in auditory object searches and were compared to listeners’ ability to detect illusory long tones, created by interspersing a 100-ms noise between two short (100-ms) tones within a cloud of other short tones and noise. Individual thresholds for perceiving the continuity illusion in the absence of other stimuli (pulsation thresholds) were also measured.

Results
Detection of physical long and short tones confirmed previously reported perceptual asymmetry based on tone length. Illusory long tones were more readily detected than the short tones, but produced statistically significantly poorer performance than the physical long tones. Individual pulsation thresholds for the illusion alone were not found to be related to ability to detect the illusion within a cloud of masking tones and noise.

Conclusion
The results suggest that the continuity illusion can occur to some extent in the presence of a complex acoustic background, and in the absence of focused auditory attention towards the target tone. Previous reports have attributed perceptual asymmetry of long and short tones to the relatively greater number of duration-specific midbrain and cortical neurons tuned to long tones. If so, our results suggest that the illusory long tones may also activate some proportion of the neurons tuned to long-duration stimuli, suggesting a relatively early, and possibly pre-attentive locus for the continuity illusion. [Supported by NIH grant R01 DC 07657.]

Spectral Motion Contrast: Another Factor in Speech Context Effects?
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Background
Spectral contrast effects, such as the auditory negative afterimage, may help normalize the incoming sound and produce perceptual constancy in the face of the variable acoustics produced by different rooms, talkers, and backgrounds. Such contrast effects may be particularly important for speech perception; indeed, a number of speech context effects have been explained in terms of static, or long-term, spectral contrasts. The present study examines another candidate for speech context effects, based on dynamic, rather than static, spectral contrasts. In particular, we attempt to dissociate effects of the long-term spectrum from those due to spectral motion or glides.

Methods
In the first experiment, we measured the effect of a preceding 500-ms spectral glide on the perceptual boundary between upward and downward spectral motion of a brief (50-ms) target glide. In the second experiment, the perceptual boundaries between [ba-da] and [da-ga] were measured for various precursors that included static and/or dynamic spectral components.

Results
Results from the first experiment confirmed earlier reports of a contrastive perceptual motion aftereffect, whereby a downward precursor glide resulted in a shift in the perceptual boundary between perceived upward and downward motion towards upward motion. The effect appeared to be spectrally local, with the strongest aftereffects occurring when the precursor and target were presented within the same spectral region. Preliminary results from the second experiment
suggest that the perceptual boundaries for phonemes can in some cases also be influenced by both the average spectral location and the spectral motion of the preceding sound.

**Conclusion**
Both static and dynamic spectral cues seem to play a role in inducing speech context effects. Thus, spectral motion contrast may be another factor that influences speech perception in natural contexts. [The first author is supported by a studentship from Advanced Bionics.]

**Acoustic amplitude-modulation detection and speech identification in fluctuating noise for cochlear implant users**

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**Background**
This psychophysical study aimed to assess whether the limited capacity of cochlear implant users to identify speech in various listening conditions (in quiet, in the presence of a steady-state or fluctuating noise) is determined by a reduced capacity to perceive slow temporal modulations.

**Methods**
This was achieved by measuring (i) phoneme identification in quiet and in the presence of a steady-state or fluctuating (8 Hz) masking noise and (ii) amplitude-modulation detection thresholds at 8 Hz for a group of 17 cochlear implant users. All patients were tested in free field at 70 dB SPL with their own clinical sound processor.

**Results**
Consistently with previous studies, modulation detection thresholds at 8 Hz were highly variable across patients, ranging from -1 to -22 dB (re: 100% modulation). Phoneme identification scores measured in quiet were correlated with modulation detection thresholds at 8 Hz. Phoneme identification scores measured with a steady-state noise masker were generally correlated with modulation detection thresholds at 8 Hz at positive signal-to-noise ratios (0 and +6 dB). Phoneme identification scores measured with a fluctuating noise masker at a negative signal-to-noise ratio (-3 dB) were not correlated with modulation detection thresholds at 8 Hz.

**Conclusion**
These results suggest that auditory sensitivity to slow temporal modulations is an important predictor of cochlear implantation success in real-life listening situations. However, sensitivity to slow modulations does not seem to explain the limited capacity of most cochlear implant patients to benefit from fluctuations in background noise maskers.
A direct test of the temporal-coherence model of auditory streaming

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Background

According to a recently proposed model of auditory scene analysis, sequences of temporally coherent sounds are grouped perceptually into a single auditory “stream”. Results consistent with this hypothesis were obtained in an earlier study, which involved an integration-promoting task. The current study sought to test the influence of synchrony on streaming using subjective measures and a segregation-promoting task.

Methods

In experiment 1, listeners indicated whether they heard sequences of alternating or synchronous tones at two frequencies (A and B) as one stream or two streams. A-B frequency separations of 1, 3, 6, 9, and 15 semitones were tested. In experiment 2, listeners detected occasional changes in the level of tones at one frequency (e.g., B) while the level of tones at the other frequency was randomly varied, thus forcing listeners to attend selectively to the target tones. In Experiment 3 the frequency of the distractor tone alternated by a small amount (1 semitone) between bursts within each trial to test whether this would facilitate segregation of the target tones.

Results

Consistent with previous findings, the results of experiment 1 showed a marked increase in the probability of reporting segregation as frequency separation increased from 1 and 9 semitones. For synchronous tones, the results were more variable: some listeners showed large effects of frequency separation while others did not; however, on average, the probability of reporting segregation for large frequency separations (6 semitones or more) was lower, and flatter, for synchronous tones than for alternating tones. Consistent with the results of experiment 1, the results of experiment 2 showed more consistent and robust increases in sensitivity (d’) with frequency separation for alternating than for synchronous tones. The results of experiment 3 showed no beneficial effect of frequency changes in the distracter tones on the segregation of the target tones.

Conclusion

Overall, the results provide partial support for models in which auditory streaming depends not just on frequency separation (e.g., channeling models) but also on the relative timing of events - with synchrony promoting stream integration, even for relatively large frequency separations. However, they also underscore the existence of large interindividual differences in the influence of relative tone timing on both “subjective” and “objective” (i.e., performance-based) measures of perceptual organization. [Work supported by grant R01 DC 07657.]

The effect of visual cues on auditory segregation

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Background

Previous work has indicated that, visual stimuli presented in synchrony with a targeted auditory stream can affect auditory segregation under certain conditions (van Ee et al., 2009). The present series of experiments are designed to elucidate the properties of this visual effect on auditory segregation and its possible mechanisms. Specifically, we aim to ascertain whether the visual effect exists in both one-stream and two-stream conditions and whether it is affected by the temporal properties of the auditory sequence.

Methods

In Experiment 1 (one-stream), acoustic sequences consisted of pure tones at 3 different frequencies, separated by 2 semitones, that were presented in random order. Subjects (N=6) were instructed to identify a ‘target pattern’ which consisted of 3 tones in a particular order. The tones were presented isochronously (REG condition) at 2 or 4 Hz, or in a random temporal pattern (RAND condition) with a mean rate of 2 or 4 Hz. Visual stimuli were a black fan shape (1/3 of a circle) with a radius of approximate 2 degrees. To reduce after-image effects, the stimuli cycled between 3 orientations (0-120 degrees, 120-240 degrees, 240-360 degrees). Visual stimulation varied according to 3 conditions: (1) no visual stimulation, (2) synchronous with the auditory signals, and (3) presented independently of the auditory signals at a rate of 3 Hz.

In Experiment 2 (two-stream), the auditory stimuli contained two sequences separated by 5 semitones, and participants were cued to attend to one of them. The temporal regularity pattern of the attended and ignored sequences was manipulated independently, resulting in 4 conditions: RAND-RAND, RAND-REG, REG-REG, and REG-RAND. Only rate
of 2 Hz was used in this experiment (experiments using faster rates are ongoing). Visual stimuli were absent or presented in synchrony with the attended auditory sequence, with the ignored sequence, or at a fixed rate of 3 Hz (independently of the auditory stimulation).

**Results**

In Exp 1, the visual stimulation had no effect on auditory pattern recognition. In contrast, in the RAND-REG and REG-RAND conditions in Exp 2, participants’ performance was improved when visual stimuli were synchronized with the attended auditory stream. Moreover, in the REG-RAND condition, visual stimuli that were synchronized with the distractor auditory stream resulted in degraded performance.

**Conclusion**

Our results suggest a role for audio-visual coherence in auditory segregation. Relevant control conditions are in progress.
might coincide with a dip or a peak in the masker envelope, increasing the signal duration would tend to increase the maximum SNR. This factor was expected to confer particular benefit in comodulated maskers due to the ability to 'listen in the dips.'

Methods
Temporal integration was assessed in normal-hearing adults in both the narrowband and the band-widening CMR paradigms. In some conditions, the signal was temporally centered in a masker envelope dip, and in others the signal was presented randomly with respect to the masker envelope.

Results
On average, temporal integration in a comodulated masker was substantially smaller when the brief signal coincided with a masker envelope dip than when it was presented at a random point in the masker envelope. For the slowest modulation rate tested (13 Hz), integration in the condition where the brief signal coincided with a masker envelope dip was less than that observed in steady noise, a result that is consistent with the integration of 'multiple looks' at the signal during epochs of improved SNR. In contrast to results for the comodulated masker, integration was insensitive to signal timing for the narrowband and random maskers.

Conclusion
The results obtained here are consistent with the conclusion that the relatively large temporal integration sometimes obtained for signals in a comodulated masker is due to greater maximum SNR for longer signals. When this factor is controlled for, integration may be comparable to or less than that observed in a steady masker.

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“Glimpsing” a target harmonic complex in a temporally interrupted masker
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Background
When two sounds (a target and a masker) are presented simultaneously, listeners are typically more successful in identifying features in the target sound if the masker sound contains temporal gaps. This benefit could be a consequence of the improved target-to-masker ratio (TMR) during the temporal gaps, or it might be interpreted that the temporal gaps in the masker provide a cue that promotes the perceptual segregation of the two sounds.

Methods
In the current study, sensitivity to changes in spectral profile was measured for a harmonic complex target sound. The target was always presented with a simultaneous harmonic masker that was temporally interrupted. Performance thresholds were collected as functions of differences in fundamental frequency (from 0 to 10 semitones) and target-to-masker ratio (from -20 to 10 dB). Besides thresholds, relative decision weights were estimated independently for the masker, the portions of the target that temporally overlapped the masker (target-overlap), and the portions of the target that were in the masker gaps (target-gap).

Results
Profile analysis thresholds improved with increasing fundamental-frequency difference, TMR, and gap duration. At low TMR’s, the target-gap stimulus dominated the responses whereas the target-overlap contributed little to the responses, indicating that listeners conducted the task by glimpsing the target in the masker gaps. At high target-to-masker ratios, the target-overlap and target-gap contributed equally to the responses.

Conclusion
When it is advantageous to do so (e.g., at low TMR’s), listeners perform profile analysis of the target using glimpses of the target during the temporal gaps of the concurrent masker. When the gap duration is short and the fundamental-frequency difference between the target and masker is small, the masker cannot be totally ignored. The efficiency of glimpsing the target improved as the fundamental-frequency difference increases.
Efficient parameter-based estimates of auditory filters

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Background
Performance-based methods currently used to estimate auditory filter parameters are time-consuming, making routine estimates difficult. An adaptive “parameter-based” Bayesian adaptive procedure for the rapid estimation of auditory filter parameters was proposed and tested.

Methods
The power spectrum model of masking was assumed with an efficiency parameter K and a two-parameter roex (r, p) filter, yielding a 3-D space of model parameters. A logistic psychometric function linked the model to behavioral responses. The task was the detection of a tone added to a notched noise, yielding a 2-D stimulus space composed of potential normalized notch bandwidths and signal strengths in dB SPL. Following Kontsevich and Tyler [1999; Vis. Res. 39, 2729-2737], a one-step-ahead search algorithm with an entropy-based criterion was adopted in which the next stimulus to be tested was the one that decreased the total expected entropy in the parameter estimation.

Results
The procedure was successful for most young normal-hearing listeners in that parameter estimates obtained from the proposed procedure and 100-150 trials were as reliable as those obtained using 2000 trials with a performance-based procedure. For a few listeners, the procedure failed to converge to reasonable parameter estimates, and for them, the performance-based measures were also variable. By introducing a “lapse” term into the psychometric function, the parameter-based procedure converged rapidly (100-150 trials) and reliably even for these few listeners.

Conclusion
The current data provided a proof-of-concept that estimates of auditory filters for young, normal hearing listeners could be achieved in as few as 150 trials. This parameter-based procedure can be extended to include other populations and other well-modeled psychophysical function.
Effects of Prior Frequency Separation and Prior Perception on Auditory Stream Segregation: Same or Different Effects?

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Background
Previous studies have shown that the perceptual segregation of low- (A) and high- (B) frequency tones in a repeating ABA pattern depended on two aspects of the previous context. First, sequences were more likely to be organized into separate streams when preceded by a context sequence that had a small frequency separation (\(\Delta f\)) between A and B tones, despite the fact that small frequency separations during the context tends to result in perception of one integrated stream during the context. Second, studies also showed that sequences with the same \(\Delta f\) as the preceding context sequence were more likely to again be organized into separate streams when preceded by a context that was also heard as segregated. Despite differences in the direction of these effects (i.e., contrastive vs. facilitative for effects of prior \(\Delta f\) and prior perception, respectively), it could be that the effect of prior \(\Delta f\) actually reflected a contrastive effect of prior perception when the \(\Delta f\) between context and test changed, and was not due to the size of the \(\Delta f\) per se. To test this hypothesis, the effect of prior perception was measured separately for conditions in which the context and test sequence had matching or mismatching \(\Delta f\).

Methods
Stimuli consisted of a repeating ABA pattern arranged into a 6.72-sec context sequence and a subsequent 6.72-sec test sequence. The \(\Delta f\) between A and B tones in the context sequence was 3, 5, or 7 semitones. The \(\Delta f\) in the test sequence was always 5 semitones. Participants continuously reported whether they heard the A and B tones grouped into the same or separate streams.

Results
Context sequences were more likely to be organized into separate streams when the \(\Delta f\) was large. Test sequences were more likely to be organized into separate streams when the preceding context \(\Delta f\) was small or was also organized into separate streams. Importantly, the effect of prior perception was facilitative regardless of whether the \(\Delta f\) between context and test matched or mismatched.

Conclusion
These results do not support the theory that the apparent effect of prior \(\Delta f\) reflected a contrastive effect of prior perception when the \(\Delta f\) between context and test changed. Therefore, we conclude that the effects of prior perception and prior \(\Delta f\) reflect distinct contextual influences of immediate prior listening. [Supported by NSF BCS1026023]

The Role of Object Perception in Change Deafness: Behavioral and Event-Related Brain Potential Findings
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Background
Change deafness is the failure to notice obvious changes occurring in an auditory scene. We sought to determine if change deafness is a fundamental sensory process, rather than a reflection of verbal memory limitations that would be predicted to only occur for recognizable sound objects. We also examined whether successful encoding of objects within a scene is related to successful detection of changes.

Methods
Event-related potentials (ERPs) were recorded while listeners completed a change-detection task and an object-encoding task with scenes composed of recognizable or unrecognizable sounds. For the change-detection task, listeners heard a scene composed of four concurrent sound objects, followed by 350 ms of silence, followed by another four-object scene that was either the same or different (three of the same objects plus one new object). Listeners then indicated whether the two scenes were the “same” or “different”. Finally, for the object-encoding task, listeners heard a probe sound and indicated whether it was present or absent in one or both of the preceding scenes.

Results
More change deafness occurred for the unrecognizable sounds, compared to recognizable sounds. ERPs from Scene 1 revealed an enhanced positivity from 315-660 ms at right temporal electrodes for detected changes, but only for recognizable sounds. ERPs from Scene 2 revealed an enhanced P3 during detected changes for both recognizable and...
unrecognizable sounds. Performance on the object-encoding task revealed that change deafness was reduced, but not eliminated, when recognizable objects were accurately encoded. Neural responses recorded during the change-detection task with recognizable sounds revealed an enhanced P2 in both Scene 1 and Scene 2 when objects were not accurately encoded.

Conclusion
The data are consistent with the conclusion that change deafness is at least partly an auditory sensory phenomenon and not solely a product of verbal memory. Enhanced neural activity for recognizable sounds in Scene 1 might reflect more accurate mapping of objects onto pre-existing representations in memory during Scene 1 or better segregation of objects based on their acoustics. The enhanced P3 for detected changes in Scene 2 may indicate that conscious change detection has occurred. A parsimonious interpretation of the finding of a smaller P2 during successful object encoding is that better performance on the object-encoding task is supported by more efficient object representations, which also leads to better change detection performance. [Supported by Army Research Office grant W911NF-I2-I-0256]

Comodulation masking release (CMR) in treefrogs and a frog-inspired auditory filterbank to investigate its neural correlates

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Background
Noisy social settings can impair the auditory system’s ability to detect, recognize and localize sound sources. Female Cope’s gray treefrogs (Hyla chrysoscelis) encounter this situation when attempting to find potential mates that call in loud, mixed-species choruses. Natural sources of noise may exhibit temporal envelope fluctuations that are correlated across the frequency spectrum, and this comodulation may be exploited by the auditory system for improved signal processing. Comodulation masking release (CMR) may depend on across-channel or within-channel frequency processing, or both. The anuran auditory periphery provides a unique opportunity to assess independently these contributions because anurans possess two distinct sensory organs in their inner ears, the amphibian papilla (AP) and the basilar papilla (BP), each responsible for processing mostly non-overlapping frequency ranges.

Methods
In this study we developed a frog-inspired auditory filterbank to investigate the correlation structure of natural frog choruses. This filterbank is more biologically pertinent than models with equal bandwidth and evenly spaced filter bands, or models based on the human auditory periphery. In a meta-analysis, a rounded exponential function (roexp) was applied to fit VIIIth nerve frequency tuning curves from several anuran species. The roexp function successfully described the filter shapes of AP and BP units, and was used to construct a filterbank.

Results
The output of this filterbank showed envelope fluctuations from different frequency bands of the chorus structure to be highly correlated. In behavioral experiments, we tested for CMR in H. chrysoscelis. Signal recognition thresholds were estimated for calls in quiet, or in the presence of “chorus-shaped noise” comprised of two narrowband noises centered on the two spectral peaks (1.3 and 2.6 kHz) present in the male advertisement call. The two narrowband noises were either 1) unmodulated (flat), 2) independently modulated (deviant), or 3) comodulated with identical modulators. At higher masker levels (73 dB SPL), subjects experienced an approximate 3 dB and 6 dB release from masking in comodulated noise compared to deviant or flat noise, respectively. This masking release was absent at a lower masker level of 53 dB SPL.

Conclusion
Our results confirm CMR-like processes that may contribute to improved call recognition in H. chrysoscelis, and we provide a first description of an anuran auditory filterbank as an analytical tool for predicting and explaining empirical results underlying mechanisms of auditory processing.

[This work supported by NIDCD R01 5R01DC009582.]
Spatial Release from Masking Improves Temporal Pattern Recognition in Cope’s Gray Treefrog
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Background
For social species, such as humans, birds and frogs, that gather in large and noisy social groups, auditory mechanisms that improve acoustic signal recognition in noise are particularly important for successful communication. One way receivers in such environments might isolate target signals from competing sounds is by exploiting spatial separation between sources. We tested the hypothesis that spatial release from masking improves temporal sound pattern recognition in Cope’s gray treefrogs (Hyla chrysoscelis). Like many other frogs, male gray treefrogs form loud breeding “choruses” where they produce calls with species-specific temporal structures that females use to recognize potential mates. To choose a mate of the correct species, females must correctly recognize the temporal pattern conveyed by the pulse rate (~50 Hz) of calls amid competing calls and high levels of background noise. This ability is necessary to avoid mating with a closely-related species with spectrally similar, pulsatile calls having a slower pulse rate (~20 Hz). If spatial release from masking improves temporal pattern recognition, we predicted females would be more likely to choose calls with faster pulse rates in the presence of artificial chorus noise originating from spatially separated sources compared with co-localized conditions.

Methods
To measure behavioral decisions of subjects, we used a phonotaxis assay requiring females to approach one of two alternating synthetic calls differing only in pulse rate. Subjects were presented with all pairwise combinations of alternatives having pulse rates of 20, 30, 40, and 50 Hz. These tests were replicated in quiet and in the presence of artificial chorus noise presented from separated or co-localized conditions.

Results
In quiet conditions, females showed strong and consistent selectivity favoring call alternatives with faster versus slower pulse rates. Hence, faster pulse rates constituted the “correct” behavioral response measured in quiet conditions. While error rates were higher in noise, subjects were significantly more likely to choose correct pulse rates when signals and noise were separated by 180° compared with the co-localized condition.

Conclusion
Spatial separation between sources improved the ability of subjects to make correct behavioral decisions based on analyses of temporal sound patterns. These results support the hypothesis that spatial release from masking improves temporal sound pattern recognition in the context of a cocktail-party-like communication problem in frogs. [This work was supported by 5R01DC009582.]
Phonemic Restoration: Studying the Effect of Voice Alternation

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Background
This study focuses on the importance of voice continuity for phonemic restoration (PR). The restoration effect can be observed by the increase in intelligibility when silent interruptions in a sentence are filled with noise, and is a measure of the auditory system’s top-down compensation for speech degradation. Poor PR performance was previously observed with cochlear-implant (CI) users, where the fundamental frequency (F0) is not well delivered and thus pitch cues are missing. Based on this, we hypothesized that the continuity of pitch and vocal-tract-length (VTL) is important for PR, as these cues can be used to group speech segments in interrupted speech. Disrupting this continuity should therefore decrease both overall intelligibility of interrupted speech and restoration effect.

Methods
Sentences originally spoken by a male talker were processed in STRAIGHT into four voice conditions: (1) unmodified vocal characteristics, (2) only F0 shifted up toward that of a typical woman value, (3) only VTL shortened to that of a typical woman size, and (4) both F0 and VTL modified simultaneously. The continuity of these vocal characteristics was then disrupted by alternating the synthesised voices across segments of meaningful interrupted sentences. Intelligibility of processed sentences was assessed with normal hearing, Dutch native speakers.

Results
Our preliminary results showed that disrupting the continuity of vocal-characteristics reduces overall intelligibility when VTL was manipulated (alone or together with F0), but not when only F0 was manipulated. However, phonemic restoration was present and unaffected in all voice conditions.

Conclusion
Although phonemic restoration is generally linked to the continuity illusion, the effect of restoration was preserved even when the continuity of vocal characteristics was disrupted to the degree that the listeners attributed different segments of speech to different talkers. In contrast, overall intelligibility was affected by the manipulation, in agreement with literature on speaker-normalization on the one hand and fusion of segments of interrupted speech on the other hand. However, the listeners perhaps took advantage of speech redundancy for restoration and compensated for distorted voice characteristics by taking advantage of other cues (such as linguistic and contextual cues). Further, as speech redundancy is very low in CI simulations and actual CIs, these results may suggest that the reduced PR effect observed in CI users or normally hearing listeners attending to CI simulations is unlikely to be only due to the poor pitch representation provided by the implant or the simulation.

Contribution of spectral and temporal cues to phonetic discrimination in infants

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1Université Paris Descartes-CNRS, 2Ecole Normale Superieure

Background
A previous study (Cabrera, Bertoncini & Lorenzi, 2011) investigated the perception of spectral and temporal cues in normal-hearing (NH) 6-month-old infants by using noise-vocoded speech stimuli and a headturning preference procedure. Noise-vocoders degraded selectively frequency-modulation (FM, the rapid variations with a rate close to the center frequency of the auditory filter) and amplitude-modulation (AM, the relatively slow variations in amplitude over time) cues. Vocoders also degraded the spectral resolution of speech signals by extracting AM within 4 broad frequency bands. After one or two minutes of familiarization, infants discriminated voicing (/aba/ versus /apa/) in each vocoded condition, but their reaction differed according to the type of degradation (see figure 1). These results suggested that the relatively fast FM and AM cues are not essential for infants to discriminate voicing in silence. The present research extended this initial study by assessing the ability of infants to discriminate two phonetic contrasts [voicing and place of articulation (/aba/ versus /apa/)] using an infant-controlled familiarization time. Tone-excited vocoders were used in order to limit potential distortions of AM cues introduced by the vocoder processing (Kates, 2011).

Methods
Four tone-excited vocoders were designed to reduce (i) FM cues, (ii) AM cues to their slowest variations (<16 Hz), (iii) and the spectral resolution of speech signals (from 32 to 8 bands). Discrimination abilities were assessed for 160 French 6-
months who reached a familiarization criterion and completed a test phase including presentation of novel and familiar stimuli.

Results
Six-month-olds discriminated voicing and place in each vocoded condition. However, differences appeared in the time required to reach the familiarization criterion: infants required more time to be fully familiarized when fast AM cues were severely reduced (see figure 2).

Conclusion
NH 6-month-olds did not require fast FM and AM speech cues to distinguish voicing and place of articulation. This corroborates the idea that fine spectro-temporal details do not play a major role in phonetic discrimination in quiet for both infants and adults. Nevertheless, longer exposure to vocoded stimuli was required when fast AM cues were attenuated, suggesting that periodic, F0-related, envelope cues are perceived and used by infants.
Effect of Temporal Characteristics of Priming on Speech Recognition
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Background

When a listener hears speech that has been degraded, a striking change in perception takes place once s/he knows the content of the message. For example, a listener first hearing speech processed through a noise vocoder with a small number of channels will hear the signal as a noisy distorted sound. This sound may or may not be interpreted as speech, and is likely to have limited intelligibility. However, after listeners are informed of the content of the message and the presentation of the vocoded sample is repeated, listeners are often impressed about how easy it is to understand the distorted speech. Quantification of the effect is difficult, because once the content is known, the listener could simply remember it and repeat it whether or not the degraded message was correctly perceived. In this study, a forced-choice discrimination task was used to alleviate this difficulty. Listeners reported whether a printed sentence (the prime) and an acoustically presented sentence were the same or different. The specific goal of this study was to determine the importance of the relative timing of the prime and the acoustic presentation.

Methods

Normal-hearing subjects listened to nonsense sentences that had been processed with a four-channel noise vocoder. They also read the sentences printed on a computer screen. On half the trials one of three key words in the nonsense sentence was replaced with a non-rhyming foil word. The subjects’ task was to determine whether the sentence they heard was the same as the sentence they read. The printed (prime) sentence was delivered at a time that ranged from two seconds before to two seconds after the acoustic presentation.

Results

As expected from previous results from a similar task using masked speech, the delivery of the written prime before the acoustic sentence produced significantly better discrimination performance than the reverse. Additional analysis focused on the comparison between the effects of simultaneous delivery of the prime versus prior delivery of the prime, and investigated the effects of fine differences in timing of the prime within a few hundred milliseconds of the acoustic presentation.

Conclusion

The effect of priming in improving speech recognition was reaffirmed. If data analysis indicates that the precise timing of the delivery of the prime is of importance, it could have implications for how priming should be delivered as part of an auditory training program. [Work supported by NIH DC01625.]
The Relative Roles of Temporal Neural Cues in Predicting Speech Intelligibility

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¹McMaster University

Background

Speech intelligibility predictors based on the contributions of envelope (ENV) and time fine structure (TFS) neural cues have many potential applications in hearing and communications research. However, establishing robust correlates between subjective speech perception and these neural cues has not been straightforward. The spectro-temporal modulation index (STMI) metric¹ has been able to predict the effects of presentation level on intelligibility in normal-hearing and hearing-impaired listeners² and the effects of different hearing aid compression schemes³. Despite these results, the STMI metric cannot explain speech intelligibility for “auditory chimaeras”⁴ where speech information is primarily in the TFS, motivating the inclusion of TFS neural cues in the predictive models.

Methods

A speech corpus of 1,750 sentences divided into five chimaera types, subjectively evaluated by 5 normal hearing listeners, was used⁵. Using sentence pairs consisting of an unprocessed NU-6 sentence and one of five respective chimaeric forms, model auditory nerve responses⁶ were simulated. The resulting spectro-temporal characterizations, termed auditory nerve “neurograms,” were subsequently processed by the STMI metric as well as the neural similarity (NSIM) metric⁶. The latter metric can capture TFS cues along with ENV related information, unlike the STMI⁵.

To investigate the ability of these metrics to predict percent-correct phoneme scores for the NU-6 chimaeras, 3 linear regression models were considered. Model 1 uses ENV and TFS with one interaction term, while Model 2 uses STMI with TFS and one interaction term. Model 3, using only the STMI metric, provides a basis of comparison.

Results

Table 1 summarizes the predictive performance of the 3 models. Relative to the STMI only, which considers only ENV cues, both Model 1 and Model 2 provide better predictive performance with the inclusion of the NSIM TFS term. It is interesting to note the combination of STMI with NSIM TFS explained more variation than the NSIM TFS and its native ENV pairing.

Conclusion

In this experimental framework, speech intelligibility predictions based solely on ENV neural cues are inadequate. Including complementary TFS cues provides better predictive performance.

¹Elhilali et al., Speech Comm 2003; ²Zilany& Bruce, IEEE EMBS Conf on Neural Eng 2007; ³Leung & Bruce, IHCON 2008; ⁴Smith et al., Nature 2002; ⁵Ibrahim & Bruce, ARO 2011; ⁶Hines & Harte, Speech Comm 2012; ⁷Zilany et al., JASA 2009

NSERC Discovery Grant 261736
Prediction of RAU Transformed % Correct CVC NU-6 Words

<table>
<thead>
<tr>
<th>Model 1</th>
<th>Model 2</th>
<th>Model 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>ENV</td>
<td>258.2 (0.0002)</td>
<td>STMI</td>
</tr>
<tr>
<td>TFS</td>
<td>-2.5 (0.986)</td>
<td>TFS</td>
</tr>
<tr>
<td>ENV x TFS</td>
<td>397.8 (0.443)</td>
<td>STMI x TFS</td>
</tr>
<tr>
<td>Adj. $R^2$</td>
<td>0.65</td>
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<tr>
<td>p-value</td>
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<tr>
<td>Adj. $R^2$</td>
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<td>Adj. $R^2$</td>
</tr>
<tr>
<td>p-value</td>
<td>&lt; 0.0001</td>
<td>p-value</td>
</tr>
</tbody>
</table>

Regression coefficients for NSIM ENV, NSIM TFS and STMI terms and their interaction are shown as in parenthesis. Overall goodness of fit as indicated by the Adj. $R^2$ value is also shown.
Consonant Identification using Temporal-Fine Structure and Recovered Envelope Cues
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Background
The present study examined the contribution of recovered envelopes (RENVs) to the utilization of temporal-fine structure (TFS) speech cues for normal-hearing (NH) listeners. The Hilbert Transform has been used to separate speech into envelope and fine-structure components; however, this processing may not completely isolate these two components. Specifically, it has been shown that when a broadband signal is filtered through a narrow filter (such as a cochlear filter of a NH listener), the TFS component yields an envelope at the output of the filter that is similar to the envelope of the band (Ghitza, 2001). Thus, speech-reception performance with TFS signals may depend on the use of RENVs in addition to rapidly-varying fine-structure cues.

Methods
Consonant-identification experiments were conducted using speech stimuli (16 consonants in /a/-C-/a/ syllables) processed to present TFS or RENV cues. For TFS conditions: speech was filtered into $N$ adjacent frequency bands over a range of 80-8020 Hz; the TFS component was extracted within each band; and the $N$ resulting TFS signals were recombined to create the TFS stimulus. For RENV conditions: the TFS stimulus was filtered into $M$ adjacent bands; the envelope components were extracted to yield the RENVs for each band; and the resulting RENVs were used to modulate tone carriers equal to the center frequency of each of the $M$ bands.

Experiment 1 examined the effects of exposure and presentation order using 16-band TFS and 40-band RENV speech. Experiment 2 examined the effect of varying the number of RENV bands from 4 to 40 with constant, 16-band TFS processing. Experiment 3 examined the effect of varying the number of TFS bands from 1 to 40 with constant, 40-band RENV processing.

Results
Preliminary results for Experiment 1 indicate that, with sufficient exposure, performance on TFS and RENV speech was comparable at roughly 55%-correct and that prior exposure to TFS speech aided in the identification of RENV speech. For Experiments 2 and 3, the dependence of performance on the number of TFS and RENV bands will be described.

Conclusion
This study investigates whether performance observed in previous studies of TFS speech may have been affected by the ability of NH listeners to recover envelopes from the TFS stimuli. The poor performance previously reported for hearing-impaired listeners with TFS speech may be related to an inability to recover envelopes due to broadened auditory filters.
The relationship between background noise envelope power and speech intelligibility in adverse conditions

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Background
Classical models of speech perception, such as in the speech intelligibility index (SII), assume that the long-term speech-to-noise ratio and the spectral characteristics of the background noise determine intelligibility. In addition, the integrity of speech envelope fluctuations has been known to play a role in conditions with reverberation. However, recent studies have demonstrated that the inherent envelope fluctuations of the background noise also play a crucial role for the speech intelligibility in adverse conditions (Jørgensen and Dau, JASA 2011; Stone et al., JASA 2011, 2012). This is reflected in the speech-based envelope power spectrum model (sEPSM) proposed by Jørgensen and Dau (JASA, 2011), which uses the signal-to-noise envelope power ratio ($SNR_{env}$) as the main predictor of speech intelligibility. Here, a multi-resolution sEPSM (mr-sEPSM) is presented and evaluated in conditions with stationary and fluctuating interferers as well as in conditions where the background noise envelope power is varied systematically using a modulation processing tool (Decorsière et al., submitted). The central hypothesis is that the speech intelligibility is a monotonic function of $SNR_{env}$.

Methods
Model predictions were compared to literature and own data for speech mixed with three stationary and five fluctuating interferers. In addition, predictions were compared to new data obtained for speech mixed with a modified stationary speech-shaped noise (SSN) where the noise envelope power was attenuated or amplified in the modulation-frequency range between 4 and 20 Hz, while keeping the overall energy of the signal the same.

Results
Lower speech reception thresholds (SRT), i.e. better speech intelligibility, were obtained with the fluctuating noises in comparison to the stationary SSN, demonstrating speech masking release (MR). When the SSN envelope power was attenuated by 20 dB, the SRT decreased by 2 dB relative to the unprocessed SSN, suggesting a release from modulation masking. When the SSN envelope power was amplified by 20 dB, the noise took on a fluctuating character and led to an MR of 10 dB. The mr-sEPSM accounted well for the data in all conditions.

Conclusion
The envelope power of the noise alone plays a crucial role for speech intelligibility in adverse conditions. The mr-sEPSM-predictions were in good agreement with the data obtained in the various experimental conditions with stationary and non-stationary interferers, supporting the hypothesis that the $SNR_{env}$ is a powerful objective metric for speech intelligibility prediction.
Abnormal Speech Processing for Hearing-Impaired Listeners in Frequency Regions where Absolute Thresholds are Normal

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Background
Speech intelligibility is reduced for listeners with sensorineural hearing loss, especially in noise. The extent to which this reduction is due to reduced audibility or supra-threshold deficits is still debated. The specific influence of supra-threshold deficits on speech intelligibility was investigated.

Methods
The intelligibility of nonsense speech signals in quiet and in various noise maskers was measured for normal-hearing (NH) and hearing-impaired (HI) listeners with hearing loss in high-frequency regions. Participants in both groups had a wide range of ages. The effect of audibility was limited by filtering the signals into low (≤1.5 kHz), mid (1-3 kHz) and low+mid (≤3 kHz) frequency regions, where pure-tone sensitivity was normal or near-normal for the HI listeners. The influence of impaired frequency selectivity on speech intelligibility was investigated for NH listeners by simulating broadening of the auditory filters using a spectral-smearing algorithm. Temporal fine structure sensitivity was estimated for NH and HI listeners by measuring sensitivity to interaural phase differences. Otoacoustic emissions and brainstem electrical responses were measured.

Results
HI listeners showed mild to severe intelligibility deficits for speech in quiet and in noise. Similar deficits were obtained for steady and fluctuating noises. Simulated reduced frequency selectivity also led to deficits in intelligibility for speech in quiet and in noise, but these were not large enough to explain the deficits found for the HI listeners. The results suggest that speech deficits for the HI listeners may result from suprathreshold auditory deficits caused by outer hair cell damage and by factors associated with aging. The influence of temporal fine structure sensitivity remains unclear.

Conclusion
Speech intelligibility can be strongly influenced by supra-threshold auditory deficits. Audiometric thresholds within the “normal” range (better than 20 dB HL) do not imply normal auditory function.
A Blind Binaural Speech Intelligibility Prediction Model
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Background
The ability to hear out speech in noisy environments is considerably enhanced using both ears compared with purely monaural listening. Brain mechanisms underlying enhanced speech recognition in noise have been studied intensively and a number of binaural models have been proposed in order to predict speech intelligibility. Despite several of these models being able to achieve high correlations with subjective ratings, however, their practical usage is limited, as they all require a priori knowledge of the room acoustics, such as the room impulse response (RIR). Here, we develop and evaluate a blind measure to predict speech intelligibility based on an auditory-inspired model.

Methods
The proposed binaural model implements an equalization-cancellation stage together with a modulation frequency estimation stage. The binaural advantage was obtained as a combination of the Better Ear and Binaural Unmasking advantages. Our model "BiSIM", was first tested in anechoic using a single speech interferer paradigm. In a second set of experiments, the model was tested in conditions involving up to three simultaneous interferers in anechoic environments.

Results
Predictions from the model were compared with those obtained with an established "non-blind" model proposed by Lavandier and Culling (2010, J. Acoust. Soc. Am. 127, 387-399), as well as with data from studies investigating subjective SRTs (Speech Receptive Thresholds). Our experimental data indicate that the proposed blind model provides reliable predictions in anechoic rooms and single interferer, with correlations as high as 0.99. In multiple masker conditions, the models can still reliably predict subjective SRTs, although smaller correlations were found.

Conclusion
The proposed binaural blind model was successfully tested in anechoic environments, and in conditions involving multiple interferers. The data also suggest the importance of modulation energies in predicting binaural speech intelligibility. Further investigations will be directed towards solving some of the model limitations.
Effects of glottal phase on negative level effects ("rollover") in speech perception
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Background
Speech recognition performance worsens above moderate presentation levels (~72 dB SPL). This "rollover" in performance is most evident in noisy conditions and is seen for both normal hearing and hearing impaired listeners. In a number of psychophysical tasks, level differences appear to influence auditory processing more for very "peaky" waveforms (such as a glottal wave) than for less modulated signals. The current research examined whether varying the shape of the glottal source exciting the speech envelope might influence the amount of rollover in speech scores at high levels.

Methods
Experienced normally hearing listeners were tested on sentences from the Modified Rhyme Test (MRT) at a +12 dB signal-to-noise ratio. Sentences were set to a levels of 48, 56, 64, 72, 80 or 88 dB SPL and then high-pass filtered with a 2.5 kHz cutoff frequency. Original MRT stimuli were presented along with stimuli that were resynthesized by applying a 64-channel vocoder to an excitation signal consisting either of noise or one of three types of harmonic complexes with different phase relationships generating different amounts of waveform peakiness (random phase, Schroeder-phase, or cosine phase).

Results
Unprocessed MRT sentences produced higher scores than any of the resynthesized stimuli. Consistent with previous findings, unprocessed sentences showed rollover with a decrease in scores of ~10% as levels increased from 72 to 88 dB SPL. The processed sentences all showed similar performance at the lowest (48 dB SPL) level but varying amounts of rollover as levels increased. The stimuli synthesized using the peakiest (cosine-phase) source function showed the most rollover, decreasing by more than 20% between 56 and 88 dB SPL.

Conclusion
The greater rollover seen for the peaky cosine-phase stimuli may be related to recent results on auditory localization (Brungart and Simpson, ProcIntConfAudDisp, 2008) which showed that vertical localization accuracy decreases for peaky click-train stimuli as levels increases from 50 to 80 dB SPL. Stimuli with less peaky waveforms did not show this level dependence. The current results may indicate that the peaky glottal waveform underlying natural speech contributes to rollover at high presentation levels. Development of speech synthesis techniques using less peaky glottal sources may lead to more intelligible speech stimuli when high presentation levels are required, as is often necessary to address audibility issues in hearing-impaired listeners.
Pilot Experiments for a Behavioral Model to Detect Tinnitus in Guinea Pigs
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Background
To confirm the presence of tinnitus in animals, behavioral models aim at assessing the inability to detect silence. We explored whether guinea pigs can be trained to detect silence. Initially, we replicated earlier experiments, that showed how guinea pigs can be conditioned to an external sound stimulus. Subsequently, we tested whether guinea pigs could be trained to detect a silent gap in noise.

Methods
Seven male guinea pigs were trained in a shuttle box with two compartments to shuttle at the presence of a conditioned stimulus (CS). When subjects failed, an air puff (the unconditioned stimulus, US) was applied as a motivator. A training consisted of 10 daily sessions, with each session containing 20 trials. Hearing thresholds were determined using auditory brainstem responses.

Results
Guinea pigs could be conditioned to a broadband noise (BBN) burst. After 10 training sessions, most animals (5/7) showed >85% correct avoidance responses (CAR) to the CS. Additionally, animals were able to generalize between sound levels ranging from 58 dB to 88 dB, where a higher sound level providing better performance. Subsequent training of these same animals to a silent gap in noise was unsuccessful.

Conclusion
We were able to replicate and confirm the findings of Agterberg et al.: Guinea pigs could be rapidly conditioned to a BBN. In contrast, gap detection performance was much poorer, possibly due to the previous trainings to BBN. Therefore, currently ongoing experiments are aimed at conditioning naïve guinea pigs to a silent gap.

Tinnitus or Hearing Loss: What is Gap Detection Actually Measuring?
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Background
Gap-induced prepulse inhibition of the acoustic startle reflex (GPIAS) has become a popular technique for tinnitus assessment in laboratory animals. This method has been praised for its relative behavioral simplicity but challenged on its fundamental principles. The skepticism is derived from the concern that sound exposed animals might be detecting hearing loss, rather than tinnitus, caused by unilateral exposure. This question should be definitively answered before this
method will be accepted as a reliable test for tinnitus. The goal of this study was to address this question. Assuming that the tinnitus percept is centrally generated, we wanted to test whether it can be detected using the unexposed (intact) ear following unilateral sound exposure.

**Methods**
To induce tinnitus, CBA/CaJ mice were sound exposed unilaterally to a narrowband noise (116 dB) centered at 12.5 kHz for 1 hour under general anesthesia (Ketamine/Xylazine). Auditory brainstem responses were recorded before and after sound exposure to insure its effectiveness. Before and at different time points after sound exposure, GPIAS performance was monitored in every mouse for behavioral evidence of tinnitus.

**Results**
Prior to sound exposure all mice demonstrated good gap detection at all frequencies tested (8, 10, 12.5, 16, 20 kHz), with two ears intact or with either left or right ear obstructed. After sound exposure about half of exposed mice displayed consistent deficits in gap detection at or at bordering the frequencies at or bordering the exposure frequency (12.5 kHz) suggesting the presence of tinnitus. Following tinnitus confirmation with two ears unobstructed, the sound exposed ear was blocked with a silicone ear plug (Kwik-Sil WPI) and the test was repeated. In this testing condition the vast majority of animals exhibited analogous gap detection deficits as in the initial test with two ears unobstructed.

**Conclusion**
Following unilateral sound exposure, GPIAS reveals gap detection deficits using the unexposed ear. This strongly suggests that these deficits are not linked to hearing loss but represent behavioral evidence of tinnitus.
Loss of gap-induced suppression of acoustic startle in hamsters following intense sound exposure
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Background
Suppression of acoustic startle by gaps in noise has been found to be reduced following exposure to intense sound or treatment with salicylate. This is often interpreted as evidence of tinnitus, the argument being that tinnitus fills in the gap when the pitch of the tinnitus is similar to that of the background noise. Here, we investigated the use of this method to test for the induction of tinnitus in Syrian golden hamsters.

Methods
We tested: 1) the relationship between startle response amplitude and startle stimulus level (57-120 dB SPL) with no background noise, 2) the effects of adding 70-85 dB SPL background noise on the startle amplitude, and 3) the effects of 50 ms gaps in the background noise on startle using various combinations of background noise and startle stimulus levels to determine what conditions are optimal for maximizing both the degree of gap-induced startle suppression and the weakening of this suppression effect following intense sound exposure. The tests were conducted in control (n=9) and intense tone exposed (10 kHz, 115 dB SPL, 4 hrs) hamsters (n=9).

Results
We found that hamsters display vigorous startle responses that grow in amplitude with increases in startle stimulus level. Exposed animals showed no change in startle thresholds relative to those in controls but showed an enhancement of startle at high startle stimulus levels. Addition of background noise preceding the startle stimulus had little effect on startle in controls animals but had a strong suppressive effect on startle in sound exposed animals. When gaps were embedded in background noise preceding the startle stimulus, startle was further suppressed at all background noise levels and at all startle stimulus levels in control animals. In contrast, there was a marked reduction of gap-induced suppression of startle in exposed animals. Thus, weakening of gap induced suppression of startle paradoxically occurred simultaneous with a strengthening of suppression of startle by background noise. The weakening of gap induced suppression of startle was not accompanied by changes in the suppression of startle by brief pulses that preceded the startle stimulus.

Conclusion
The loss of gap suppression after sound exposure without a corresponding change in prepulse inhibition is consistent with the interpretation that the exposure caused induction of tinnitus in hamsters. The results are also suggestive of different inhibitory mechanisms that control startle magnitude which are differentially affected by acoustic overstimulation. (Supported by NIH grant R01 DC009097).
Two Major Improvements for Using the Gap Detection Method to Assess Tinnitus-like Behavior in Mice

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Background

Tinnitus is a major, growing health problem that affects the quality of life of millions of people. While formally defined as the perception of sound in the absence of a physical sound stimulus, people with tinnitus also report concentration difficulties, irritability, sleep disruption, anxiety and depression. There currently is no reliable treatment for tinnitus mainly due to a lack of understanding of its molecular mechanisms. With extensive molecular tools such as transgenic methods, mouse models for tinnitus would greatly facilitate this area of research. Although several animal behavior models including the gap detection method have been developed to study tinnitus in various types of animals, these methods have been difficult to apply in mouse studies. One major weakness of the gap detection method for mice is the use of average amplitude ratios between gap prepulse inhibition reflex and startle reflex as a measurement for tinnitus-like behavior. There are two reasons behind this weakness. First, the amplitude reduction of the startle reflex has been consistently observed after tinnitus induction. In addition, the neural pathway is different between the startle reflex and gap prepulse inhibition reflex. Second, we have found that most of behavior data is not normally distributed.

Methods

To solve these two issues, we first adopted the Kernel density method to select data for each trial, which solved the issue of abnormal data distribution. Next, we use the amplitude ratios between gap prepulse inhibition reflex and prepulse inhibition reflex as a measurement for tinnitus-like behavior.

Results

With these two major improvements, we can detect tinnitus-like behavior at specific frequencies in most of mice only one hour after sodium salicylate injection (350 mg/kg). This modified method is currently being tested in a noise-induced tinnitus mouse model.

Conclusion

One main goal of our studies is to study molecular mechanisms underlying tinnitus pathology in mice. With our two major improvements, the gap detection method can be an effective way to determine tinnitus-like behavior in mice, which paves way for us to study molecular signaling pathways involved in tinnitus.

Simulated small arms fire exposure in rats results in reduced gap inhibition of the acoustic startle reflex suggesting induction of tinnitus

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Background

Hearing loss resulting from exposure to ammunition discharge is a common finding in our military population. Both hearing loss and tinnitus are the leading causes for disability discharge from military service. Prevention or reduction of hearing loss and tinnitus would be of great social and economic benefit. Here we present preliminary findings of an animal model using simulated small arms fire (SAR) to induce hearing loss and tinnitus.

Methods

Sprague Dawley rats were unilaterally exposed to 50 biphasic impulses via a compression driver over a 2.5 minute period. The impulse level was either 152 or 160 dB SPL. Control rats received sham noise exposure. ABR measurements evaluated the effect of the exposure. The acoustic startle response (ASR), pre-pulse (PPI) and gap inhibition (GI) were used to evaluate the development of tinnitus.

Results

Sham treated animals showed stable ABR thresholds and consistent ASR, PPI and GI throughout the experiment. In general, animals exposed to the 160 dB stimulus showed significant unilateral ABR threshold shifts and reduced startle amplitudes sometimes resulting in “floor” effects and inconclusive PPI and GI ASR results. Animals exposed to the 152 dB stimulus showed reduced elevations in ABR thresholds compared to the 160 dB exposure. Startle amplitudes were either unaffected by the 152 dB exposure or increased in amplitude. PPI remained intact post exposure. In over half of
the animals receiving the 152 dB stimulus GI decreased, suggesting the presence of tinnitus. In some animals, the presence of the gap in noise caused enhancement of the startle response, suggestive of the presence of hyperacusis.

**Conclusion**
More than half of the rats exposed to SAR exhibited reduced GI of the ASR but intact PPI, indicative of the induction of tinnitus by this exposure condition.
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Effect of low-level laser therapy on salicylate-induced tinnitus in rat
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**Background**
Tinnitus is a perception of sounds without external source. Mainly because of the subjective nature of the disorder and our lack of knowledge of its pathophysiology, treatment of tinnitus has been unsuccessful. Low-level laser therapy (LLLT) has been introduced as an alternative treatment for tinnitus. Although many studies have been conducted for assessing a meaningful relief of tinnitus intensity, there has been no study to quantify the effect of LLLT on the treatment of tinnitus in animal model. Therefore, the aim of this study was to measure the effect of LLLT on salicylate-induced tinnitus in the rat model using GPIAS (gap prepulse inhibition of acoustic startle).

**Methods**
10 Spraque-Dawley rats (8 weeks; 240~280g) were divided into 2 groups, one for the laser group (n=5) and another for the control group (n=5). Rats of both group were treated with 400mg/kg/day of sodium salicylate for 8 consecutive days. Tinnitus was monitored using GPIAS 2 hours after first salicylate treatment, and every 24 hours during 9 days of treatment. Rats in laser group was irradiated with wavelength of 830 nm diode laser at 165mW/cm² for 30 min to each ear daily for 8 days.

**Results**
GPIAS value of both two groups was significantly decreased after sodium salicylate administration, which means tinnitus was well induced in both groups. The day after treatment start, LLLT had a tendency to increase GPIAS value of study group and a statistically significant difference was noted at 120 hours after drug administration. No significant changes in NBPIAS were observed before and after salicylate injection over the entire testing period.

**Conclusion**
Our result shows that LLLT may provide a feasible therapeutic approach to control tinnitus. Further experimental studies are needed to find better method to facilitate LLLT efficacy and possible mechanism.
Single Dose Alcohol has Protective Effect on Behavioral Deficits Post Blast-Exposure.

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Background
Alcohol has been shown to have both neuroprotective and deleterious effects after traumatic brain injury. Alcohol acts on numerous receptors within the brain (including GABAergic receptors). Non-chronic alcohol consumption works by keeping GABAergic receptors open longer, and thus increasing neural inhibition. During blast-exposure, we have observed micro-tears centered on GABAergic neurons. We hypothesize that single-use alcohol may increasing the efficacy of intact or slightly damaged GABAergic neurons and thus help re-stabilize circuitry that has sustained damage to GABAergic neurons.

Methods
To test this hypothesis, animals were orally gavaged ethanol either immediately prior or 2 days post blast exposure. They were monitored for motor ability with RotoRod and Versimax. They were also monitored for tinnitus with acoustic startle tests. To allow statistical comparisons of behaviors pre/post blast-exposure in each animal, all tests were performed both prior and post blast-exposure.

Results
Thirty minutes after alcohol administration all animals had a significant motor impairment. Analysis of behavioral measures pre/post blast-exposure indicated that the alcohol groups performed better in motor tests and had decreased incidences of tinnitus after blast-exposure.

Conclusion
We conclude that single-dose alcohol has protective effects on behavioral measures of injury after blast-induced traumatic brain injury.
Prevalence of tinnitus and its relationship with anxiety and depression
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Background
In the 1980s, about 10% of UK adults reported experiencing prolonged, spontaneous tinnitus lasting for more than five minutes. However, the current prevalence is unknown and large-scale studies of tinnitus are lacking. Furthermore, although many people learn to live with tinnitus, for some people it can be severely distressing. Clinical studies indicate a strong association between tinnitus and mental health, with increased risk for depression and anxiety among tinnitus patients. However little is known about how the severity of tinnitus is related to these disorders. The aim was to update the prevalence of tinnitus and further examine the association between the severity of tinnitus and anxiety and depression.

Methods
A cross-sectional design study of 502,713 people (54.4% female; 45.6% male) from across Britain aged between 40-69 years who participated in UKBiobank between 2006-2010. UKBiobank is a resource set up to support research intended to improve the prevention, diagnosis and treatment of illness. Two questions on tinnitus asked whether participants had experienced ringing or buzzing for more than five minutes at a time, and if so how much it worries, annoys or upsets them. Participants were also asked whether they had visited a GP/psychiatrist for nerves, anxiety, tension or depression, and specific questions on depression, including whether they had ever been depressed for at least a week, their longest period of depression, and number of depressive episodes. Associations between these factors and tinnitus characteristics were investigated using multiple regression techniques.

Results
17.7% of participants experienced tinnitus at least some of the time and, of these, 20.2% (3.6% of the total) had moderate or severe tinnitus, suggesting the prevalence of tinnitus in this age group could be much higher than previously thought. Of the whole sample, 34.1% indicated having seen a doctor (11.6% had seen a psychiatrist) for nerves, anxiety, tension or depression. 53.2% of the sample indicated being depressed for at least a week. The mean longest period of depression was 15.9 weeks and the mean number of depressive episodes was 6.3.

Conclusion
Data on prevalence and severity of tinnitus within the UK Biobank participants, and its relationship with anxiety and depression, suggest a stronger than expected association between these conditions in the UK’s middle-aged population. These data will contribute to appropriate resourcing, prevention, identification and management strategies for people with tinnitus in the medium term.
Prefrontal cortex activity is related to the tinnitus percept – group differences and correlations revealed by functional magnetic resonance imaging

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Background
Tinnitus, a phantom “ringing in the ears” affecting 10-15\% of the population, is commonly attributed to changes in the central auditory system following damage to the peripheral auditory system. There is also increasing evidence for a role of non-auditory (especially limbic and prefrontal) brain areas in tinnitus, e.g. from group comparisons based on anatomical magnetic resonance imaging and resting-state magnetoencephalography (MEG). Compared to MEG, functional magnetic resonance imaging (fMRI) has superior spatial resolution; however, the vicinity of the sinuses leads to signal loss in ventral prefrontal cortex. The present study used a modified data acquisition scheme that ameliorates this problem and employed multiple conditions with auditory stimulation to further elucidate the role of ventromedial prefrontal cortex (vmPFC) in tinnitus.

Methods
Functional MRI was performed on 20 tinnitus patients and 20 control participants without tinnitus, who were matched for age, sex, and hearing loss. Each patient, and his or her matched control, listened to band-passed noise bursts at center frequencies spanning the human hearing range, one of which was matched to the patient’s tinnitus frequency. In a control condition, no sound was played. To avoid compromising the conditions by scanner noise, fMRI data were acquired intermittently rather than continuously, and images were acquired in tilted slices to avoid cutting through the sinuses and improve the fMRI signal in ventral prefrontal cortex. Questionnaires assessed tinnitus-related measures such as tinnitus loudness, tinnitus distress, noise sensitivity, depression, and anxiety.

Results
When comparing patients and controls regarding the fMRI signal change between the condition with stimulation at the tinnitus frequency and the condition without stimulation, significant group differences were found in right vmPFC and right superior temporal gyrus (STG), including the putative location of primary auditory cortex. Post-hoc analyses constrained to these regions revealed a strong correlation between tinnitus loudness and vmPFC activation in tinnitus patients, especially during stimulation at the tinnitus frequency ($r = 0.74$). In contrast, correlations with tinnitus distress, noise sensitivity, depression, and anxiety were negligible.

Conclusion
These fMRI findings add to the evidence that vmPFC plays a role in tinnitus perception. The absence of correlations between vmPFC activation and measures of depression, anxiety, and tinnitus distress indicate that this role is not in the emotional evaluation of the tinnitus percept. Instead, the strong correlations between vmPFC activation and tinnitus loudness agree with our hypothesis that vmPFC activation can modulate the tinnitus percept (Rauschecker et al., 2010).

Tinnitus-Related Dissociation between Cortical and Subcortical Neural Activity in Humans with Mild to Moderate Sensorineural Hearing Loss.

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Background
Tinnitus is a phantom sound percept that is strongly associated with peripheral hearing loss. However, only a fraction of all hearing-impaired subjects develops tinnitus. This may be based on differences in the function of the brain between those subjects that develop tinnitus and those that do not.

Methods
Cortical and sub-cortical sound-evoked brain responses in 34 hearing-impaired chronic tinnitus patients and 19 hearing level-matched controls were studied using 3-T functional magnetic resonance imaging (fMRI). Auditory stimuli were presented to either the left or the right ear at levels of 30-90 dB SPL. We extracted neural activation as a function of sound intensity in eight auditory regions (left and right auditory cortices, medial geniculate bodies, inferior colliculi and cochlear nuclei), the cerebellum and a cinguloparietal region.

Results
The activation correlated positively with the stimulus intensity, and negatively with the hearing threshold. We found no differences between both groups in terms of the magnitude and lateralization of the sound-evoked responses, except for
Conclusion
Our results suggest abnormal thalamic gating in the development of tinnitus.

Functional Connectivity Between Auditory, Default-Mode, and Cognitive-Control Networks in Tinnitus and Hearing Loss
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Background
It is hypothesized that aberrantly correlated activity among brain areas may be important to sustaining the tinnitus condition, defined as the perception of phantom sound and the emotions surrounding the percept. The present study tested this hypothesis by comparing functional connectivity between human subjects with tinnitus and without. Functional connectivity indicates the degree to which activity in one brain area is temporally correlated with that in another.

Methods
Functional magnetic resonance imaging (fMRI) was performed on subjects in three groups: (1) high-frequency hearing loss and tinnitus (HFL-tinnitus), (2) clinically normal hearing sensitivity and tinnitus (cNH-tinnitus) and, (3) no tinnitus and threshold-matched to group 2 (cNH-control). A structural MRI of each subject was also obtained and segmented into regions of interest (ROI) comprising cortical gray matter. The ROIs were used to seed connectivity analyses of the fMRI data. Specifically, a correlation coefficient was calculated between the fMRI signal time-course within an ROI and the signal time-course of each voxel in the brain, resulting in a spatial map of correlations with the ROI seed. White matter and cerebrospinal fluid signals were regressed out. Maps of correlation for each subject were smoothed and entered into a second-level analysis for inter-group comparison.

Results
In the absence of any explicit sound stimulus, primary auditory cortex (PAC) of the cNH-control group showed: (1) positive correlations with both adjacent and contralateral auditory cortical areas (including contralateral PAC), (2) positive correlations with supplementary motor cortex (SMA) and pre-SMA, nodes of a previously-defined cognitive control network, (3) negative correlations with posterior cingulate/precuneus (PCC), right and left parietal lobes and, medial prefrontal cortex. The regions showing negative correlations are major nodes in the default mode network, which is active when external demands (e.g. listening) are not being placed on a subject. While cNH-tinnitus and HFL tinnitus subjects showed correlations between PAC and the same brain areas as cNH-controls, the magnitude of the correlations with non-auditory areas (but not auditory ones) was reduced. The reductions were significant in SMA/pre-SMA and PCC. They were greatest for the HFL tinnitus group.

Conclusion
The results demonstrate a conduit by which exogenous auditory information can interact with endogenous signals of the cognitive-control and default-mode networks and further suggest that this conduit may not operate normally in those with tinnitus, and especially those with hearing loss and tinnitus.
The Relation between Perception and Brain Activity in Gaze-Modulated Tinnitus

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Background
Tinnitus is a phantom sound percept that can be severely disabling. Its pathophysiology is poorly understood, partly due to the inability to objectively measure neural correlates of tinnitus. Gaze-modulated tinnitus (GMT) is a rare form of tinnitus that may arise after vestibular schwannoma removal. Subjects typically describe tinnitus in the deaf ear on the side of the surgery that can be modulated by peripheral eye gaze. This phenomenon offers a unique opportunity to study the relation between tinnitus and brain activity.

Methods
We included 18 patients with GMT and 9 controls. Pure-tone audiometry was performed in all subjects. Subjects with tinnitus also performed tinnitus matching tasks for six gaze directions. Cortical and sub-cortical brain responses were studied using 3-T functional magnetic resonance imaging (fMRI). The scanning paradigm included the same six gaze directions and one bilateral sound condition.

Results
In normal-hearing control subjects, peripheral gaze resulted in inhibition of the auditory cortex, but no detectable response in the medial geniculate body and inferior colliculus. In patients with GMT, peripheral gaze (1) reduced the cortical inhibition, (2) inhibited the medial geniculate body, and (3) activated the inferior colliculus. Furthermore, increased tinnitus loudness was represented by increased activity in the cochlear nucleus and inferior colliculus and reduced inhibition in the auditory cortex.

Conclusion
The increase of cochlear nucleus and inferior colliculus activity with peripheral gaze is consistent with models of plastic reorganization in the brainstem following vestibular schwannoma removal. The decrease of activity in the medial geniculate body and the reduced inhibition of the auditory cortex support a model that attributes tinnitus to a dysrhythmia of the thalamocortical loop, leading to hypometabolic theta activity in the medial geniculate body. Our data offer the first support of this loop hypothesis of tinnitus, independent of the initial experiments that led to its formulation.
Auditory evoked magnetic fields in individuals with tinnitus
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Background
Recent theories postulate that some forms of tinnitus are likely to be perceptual consequence of altered neural activity in the central auditory system triggered by damage to the auditory periphery. Despite reduced input from the cochlea and auditory nerve after noise exposure or ototoxic drugs, many animal studies have observed changes in sound-evoked firing rate, as well as amplitude of the evoked responses in inferior colliculus and auditory cortex. However, human electrophysiological evidence is rather equivocal. While some studies reported increased N1/N1m amplitude and shorter latencies in tinnitus patients compared to controls, others reported reduced N1/N1m amplitudes and prolonged latencies, and some found no differences.

Methods
The present study used magnetoencephalography (MEG) to seek evidence for altered evoked responses in people with tinnitus compared to controls. We have measured evoked magnetic fields in response to 300-ms tone bursts were presented at 40 dB SL in 22 people with tinnitus as well as in 13 participants with normal hearing and no tinnitus and 7 participants with matched hearing loss but without tinnitus. We applied channel-space analysis to compare the amplitudes and latencies of N1m responses to four different tone categories: a control tone, edge frequency, dominant tinnitus pitch and a tone within the area of hearing loss. All participants were screened to exclude hyperacusis.

Results
In agreement with the previous literature we found decreasing N1m amplitudes and only minimal shift in N1m latencies with increasing frequencies. Amplitudes of the evoked responses differed depending on the tone category, but there was no difference in this pattern across tinnitus and control groups. Overall, N1m amplitudes to the dominant tinnitus pitch were significantly smaller than for the control and audiometric edge frequencies and were similar to the tone within the hearing-loss region.

Conclusion
Dominant tinnitus pitch is typically within the area of hearing loss. We thus conclude that differences in the evoked responses pattern in participants with tinnitus were related more to the hearing loss than to the presence of tinnitus. This indicates lack of a generalised tinnitus-specific hyperexcitability in response to sound stimuli.
"Distressed aging" - The differences in brain activity between early- and late-onset tinnitus

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Background
Recent findings regarding different characteristics according to the age of tinnitus onset prompted us to conduct a study on the differences in tinnitus-related neural correlates between late-onset tinnitus (LOT; mean onset age, 60.4 years) and early-onset tinnitus (EOT; mean onset age, 29.7 years) groups.

Methods
We collected quantitative electroencephalography findings of 29 participants with LOT and 29 with EOT, as well as those from 59 controls. Then, LOTs were subdivided into 14 high distress (HD)/15 low distress (LD), while EOTs into 14 HD/15 LD. We then compared the results using resting state electroencephalography source-localized activity and connectivity analyses.

Results
Compared to the EOT and older control groups, the LOT group demonstrated increased localized activity and functional connectivity in components of previously described tinnitus distress networks, as well as the default mode and intrinsic alertness networks, such as the prefrontal corticies, dorsal anterior cingulate cortex, and insulae. On subgroup comparisons, the ACC was activated more in HD-LOT participants than in LD-LOT participants, while the orbitofrontal cortex was activated more in HD-EOT than in LD-EOT.

Conclusion
The current findings of intrinsic differences in tinnitus-related neural activity between the LOT and EOT groups may be applicable for planning individualized treatment modalities according to age of onset. Moreover, differences with regard to the age of tinnitus onset may be a milestone for future studies on onset-related differences in other similar pathologies, such as pain or depression.

Establishing the Tinnitus/Epilepsy Connection

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Background
Throughout history, parallels have been drawn between tinnitus and epilepsy. Tinnitus is described as the perception of ringing or buzzing in the absence of external stimulation while epilepsy is often characterized by seizures that are recurrent and unprovoked. Similarities between the two conditions include suggested mechanisms related to underlying electrophysiology (neuronal hyperactivity, bursting discharges, neural synchrony across brain areas), alterations in intrinsic neurochemistry (reduced intra-cortical inhibition), and changes in gamma aminobutyric (GABA) and glutamate receptors after kindling and noise induced hearing loss (Valentine et al., 2004). There are also treatment modalities common to both, including: pharmacological, electrical (vagus nerve stimulation, VNS), magnetic (repetitive transcranial magnetic stimulation, rTMS), and deep brain stimulation (e.g., Engineer et al., 2011; Connor et al., 2012; Fisher, 2012). Understanding the link between tinnitus and epilepsy may provide new treatment strategies for these conditions.

Methods
We have developed and collected results from an online survey made available to members of the American Tinnitus Association (ATA). The survey remained accessible for five weeks and more than 250 members with tinnitus (between the ages of 18 and 80) responded.

Results
The incidence of epilepsy has been reported to be 3% of the population in the United States. The prevalence of active epilepsy in our society has been reported to be as high as 1%. In the current study we found that five percent of the respondents that report tinnitus were also diagnosed with epilepsy.

Conclusion
These results suggest that epilepsy may occur with greater frequency in the tinnitus population than in the general population. Further investigation into a possible tinnitus/epilepsy link should include surveys regarding tinnitus perception
administered to epilepsy sufferers. These survey results begin to lay a foundation for studies examining common mechanisms between tinnitus and epilepsy.

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Defining Uncertainties in Tinnitus Treatments.
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Background
A James Lind Alliance (JLA) Tinnitus Priority Setting Partnership has identified and prioritise unanswered research questions about tinnitus assessment, diagnosis and treatment. This process gives a voice to members of the public and clinicians and is driven by them.

Methods
Our methods included the following four steps: (1) established a working partnership between representatives for patients and clinicians; (2) designed and disseminated a questionnaire to identify and collect tinnitus uncertainties; (3) reduced the 2483 submitted questions down to a list of 170 edited questions by pooling duplicates and removing questions that had already been answered by systematic review; and (4) prioritised these remaining questions to create a ‘top 10’.

Results
The highest ranked question (A) (Table 1) concerned management strategies and combined three separate questions asking about lifestyle changes, promoting habituation and learning to live with tinnitus. This is of the most concern to people with tinnitus. Systematic reviews have criticised the poor quality of trials [1-2] and an international standard has been proposed [3]. For hearing healthcare professionals, the second question (B) asked whether cognitive behaviour therapy (CBT)/psychological therapy, delivered by audiology professionals, is effective for people with tinnitus. Systematic reviews support a moderate benefit to patients from CBT [4], but studies tend to involve trained psychologists delivering the therapy which is not the usual model of care in the UK. C addressed the effectiveness of management strategies for patients with tinnitus and insomnia. Whilst sound therapy is commonly used by tinnitus patients at night, it is unknown whether it improves sleep because this is rarely an outcome measure [5]. We will present evidence to support all 10 research uncertainties and propose research strategies to take this work forward.

Conclusion
The top 10 questions are now registered on the NHS Evidence UK Database of Uncertainties about the Effects of Treatments (DUETs). It is managed by the National Institute for Health and Clinical Excellence (NICE). Outcomes are being widely disseminated to major stakeholders including tinnitus researchers and funders of biomedical research.

References:
The top 10 tinnitus research uncertainties.

| A | What management strategies are more effective than a usual model of audiological care in improving outcomes for people with tinnitus? |
| B | Is Cognitive Behaviour Therapy (CBT), delivered by audiology professionals, effective for people with tinnitus? Here comparisons might be with usual audiological care or CBT delivered by a psychologist. |
| C | What management strategies are more effective for improving tinnitus-related insomnia than a usual model of care? |
| D | Do any of the various available complementary therapies provide improved outcome for people with tinnitus compared with a usual model of care? |
| E | What type of digital hearing aid or amplification strategy provides the most effective tinnitus relief? |
| F | What is the optimal set of guidelines for assessing children with tinnitus? |
| G | How can tinnitus be effectively managed in people who are Deaf or who have a profound hearing loss? |
| H | Are there different types of tinnitus and can they be explained by different mechanisms in the ear or brain? |
| I | What is the link between tinnitus and hyperacusis (over-sensitivity to sounds)? |
| J | Which medications have proven to be effective in tinnitus management compared with placebo? |

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**Epidemiology and risk factors of 'significant' tinnitus: A UK retrospective study using national health service (NHS) records**

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**Background**

Tinnitus remains one of the commonest chronic hearing-related conditions in the Western world. Long-term incidence studies are particularly valuable for identifying factors associated with developing the condition. This knowledge can inform the allocation of public health resources or target prevention campaigns. Only two studies (Beaver Dam, Wisconsin, USA and Blue Mountains Hearing Study, Sydney, Australia) have so far assessed the long-term incidence of tinnitus in the general population. Their approach to risk factor modelling had some limitations. The aims of this study were to estimate the incidence rate of 'significant' tinnitus that burdens the NHS in the UK and to identify risks factors for developing the condition based on factors defined *a priori*.

**Methods**

Data was extracted from the Clinical Practice Research Datalink and Hospital Episode Statistics between 1 January 2001 and 31 December 2011. 'Significant' tinnitus was operationally defined as patient access to primary care and onward secondary referral, with a recording of tinnitus (n = 23,672). A gender- and age-matched control cohort included patients without a recording of tinnitus (n = 53,958). The choice of risk factors was defined according to a broad literature search (19 categories) and a breakdown according to period of exposure to various medications.

**Results**

The 10-year cumulative incidence of 'significant' tinnitus was 55.4 per 10,000 (95% CI:54.6-56.2). In other words, approximately one in 180 patients in the total population is likely to experience significant' tinnitus within a 10-year period. Incidence rate steadily increased during the study period. Risk analysis used a conditional logistic regression model which controlled for confounding bias and accounted for age and hearing loss. Relative risk estimates are presented as adjusted odds ratios and 95% CI. Many otologic conditions increased risk, as did head injury, rheumatological disease, anxiety and depression. Tinnitus incidence appeared to be unaffected by heart disease, use of lipid controlling drugs and alcohol consumption. Histories of diabetes mellitus, smoking, treated hypertension and pregnancy emerged as potential protective factors.

**Conclusion**

Tinnitus is a symptom that may be part of a specific otological condition but can also be associated with non otological disease processes. Unfortunately there is no objective measurement for tinnitus and consequently diagnostic uncertainty
exists, meaning that the NHS data should be interpreted with some degree of caution. Nevertheless these epidemiological
data support the growing case for developing an effective referral and management pathway for people with tinnitus.

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**Changes of tinnitus in sudden sensorineural hearing loss: the relationship between tinnitus pitch and the audiogram**

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**Background**

Different mechanisms of tinnitus, such as lateral inhibition and homeostatic plasticity, have been proposed. The aim of this study was to explore the changes of tinnitus in patients with SSNHL at initial and 1 month follow-up examinations by analyzing the relationship between tinnitus pitch and audiogram shape.

**Methods**

Thirty-six patients with SSNHL and new onset tone-like tinnitus were enrolled. The tinnitus pitch was compared to the frequency of maximum hearing loss (HL) and the audiogram edge at initial and 1 month follow-up examinations.

**Results**

At initial examination, the tinnitus pitch was in the same range with the edge frequency. The tinnitus pitch showed a significant correlation with both the edge frequency and the frequency of maximum HL. The edge frequency ($r=0.461$, $p=0.005$) had more significant correlation than the frequency of maximum HL ($r=0.333$, $p=0.047$). At 1 month follow-up, the tinnitus pitch increased significantly compared to the initial stage and was similar to the frequency of maximum HL. The tinnitus pitch showed a significant correlation with the frequency of maximum HL ($r=0.519$, $p=0.001$), but not with the edge frequency ($r=0.155$, $p=0.366$).

**Conclusion**

Significant relationship of tinnitus pitch with the edge frequency at initial examination suggests that reduced lateral inhibition and cortical reorganization play a role in generating tinnitus at acute stage of hearing loss. Significant relationship of tinnitus pitch with the frequency of maximum HL at initial and follow-up examinations suggests that tinnitus is generated and persists by homeostatic mechanism.

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**A new in vitro model to investigate the biological mechanisms involved in the alteration and restoration of the vestibular neuronal network**

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**Background**

Background: It is commonly assumed that a large proportion of vestibular disorders results from transient synaptic decoupling between vestibular hair cells and their cognate nerve afferents. However the poor spatial resolution of current imaging techniques available in human prevents the direct observation of the biological changes that occur following the synaptic insult. Using an animal model of vestibular excitotoxic insult, we previously demonstrated that selective impairment of the first sensory-neural synapse triggers the expression of altered vestibular behavior that replicates the symptomology encountered in human vestibular disorders. We also reported a high propensity of the vestibular synapses to spontaneously repair (Brugeaud et al. 2007). Such synaptic plasticity has been further confirmed in an organotypic culture model (Travo et al. 2012). More recently, we deciphered the time course of the vestibular lesion and functional restoration in an animal model of vestibular excitotoxic insult and demonstrated the correlation between the synapse impairment and the altered vestibular behavior (Gaboyard et al. poster # 247 ARO 2011). Present study was designed to develop an experimental model to access the intimate mechanisms of the vestibular lesion and restoration through several technical approaches such as imaging, histology, molecular biology and electrophysiology.

**Methods**

Method: this in vitro model consists in a three-dimensional mouse organotypic culture derived from previously detailed protocol (Gaboyard et al., 2005). Briefly, vestibular sensory epithelia and Scarpa’s ganglion were removed with particular attention paid to preserve synaptic connections between hair cells and primary neurons. Organotypic slices were produced and maintained in culture up to 15 days and the synaptic integrity studied using a multidisciplinary approach: immunocytochemistry, 3D reconstruction and quantification by morphometric analysis and, transmission electron
microscopy. Finally, the function of the synaptic transmission has been evaluated in parallel using the patch-clamp technique.

Results
Results: Observations of our in vitro model at different time point allowed us to show: (1) a clear integrity of most hair cells and vestibular neurons as attested by preservation of intact mitochondria and subcellular organelles; (2) a synaptic plasticity process with uncoupling and spontaneous repair occurring with a time course comparable to our previously described in vivo model (Gaboyard et al. 2011); 3) newly formed synapses in the process of maturation: incomplete calyces present synaptic ribbons with many vesicles facing synaptic post-densities.

Conclusion
Conclusion: This model provides a new tool to further investigate the intimate mechanisms of protection and/or repair by imaging, electrophysiology and molecular techniques.

Experimental Mechanical Response of the Utricle to Dynamic Inertial Stimulus
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Background
Few experimental studies have made direct measurement of otolith organ mechanical properties and none using dynamic stimuli. All previous analysis and experimental efforts regarding utricle mechanical motion have always concluded it is an overdamped system. Here, the utricle of the turtle was oscillated over a frequency range using an inertial stimulus, causing a measurable shear displacement of the combined gelatinous and column filament layers, the shear layer (SL). This experimental work revealed that the utricle is an underdamped system.

Methods
The utricle and its surrounding membranous sac were removed from the skull and maintained under compatible aqueous solution throughout. The membranous sac was opened to expose the dorsal surface of the otoconial layer (OL) of the utricular macula. The utricle periphery was then fixed to a glass slide with strands of dental floss. The glass slide was mounted onto a piezoelectric actuated platform that was located in a fixed stage light microscope. The piezoelectric actuator oscillated the neuroepithelium layer (NEL) of the utricle in its plane of normal acceleration stimulus. A controlled sinusoidal-sweep that increased from 0 to 500 Hz over 5 seconds was applied. This natural inertial stimulus produced a measurable shear displacement in the SL. The displacements of a reference point for the NEL and OL were filmed at 2000 frames/sec with a high-speed video camera during the oscillations. Image registration was performed on the video to track OL and NEL reference point displacements to better than 15 nm. The displacement waveforms were converted into frequency response data, and then used in a frequency domain system identification technique to determine the system mechanical properties of: (1) natural frequency, and (2) damping coefficient of the shear layer.

Results
Properties were measured for 21 utricles with a mean and 95% confidence interval in parenthesis: natural frequency \( \omega_n = 373 \text{ (350, 396)} \text{ Hz} \), damping ratio \( \zeta = 0.50 \text{ (0.47, 0.53)} \), and shear modulus \( G = 9.52 \text{ (8.32, 10.71)} \text{ Pa} \). Within the acceleration ranges tested the utricle behaved as a linear system, nonlinearities were not observed in amplitude or frequency.

Conclusion
The results indicate that the utricle performs as a linear undamped second-order system for inertial stimuli. In comparison to an overdamped system (\( \zeta > 1 \)), a \( \zeta = 0.50 \) provides improved transient-response to head accelerations, increasing bandwidth.

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Noise-induced vestibular deficits in rats
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Background
The vestibular system responds naturally to head acceleration. Because vestibular end organs are connected to the auditory end organs by continuous fluid pathway within the membranous labyrinth, the vestibular system also responds to sound. It is well known that extensive noise exposure leads to sensorineural hearing loss. It has been suggested that noise exposure also leads to vestibular deficits in humans, but little is known about the underlying mechanisms. In the present study, we established a unilateral noise-induced hearing loss rat model to study the influence of noise exposure on the vestibular system.

Methods
Sprague-Dawley rats were anesthetized with isoflurane. Continuous broadband white noise at an intensity of 116 dB pSPL was delivered to the left ear via an insert ear phone. A hard plastic infant tip adapter was placed at the end of sound-conducting tubing and placed into a speculum that was sealed in the ear canal. The contralateral ear canals were sealed from air-conducted sound exposure. The rats received noise exposure 3 hours per day for 5 consecutive days. Hearing level was monitored by measuring ABR threshold (0.1 ms clicks).

Results
After 5 days of exposure, ABR thresholds of the ipsilateral ears increased from 28 ±2.7 dB HL to 70.8±7.8 dB HL and were at 64.5±3.0 dB HL 7 days post-noise exposure. A 20-30dB temporary threshold shift was observed on the contralateral side, which may result from bone-conduced noise exposure from the left side. The vestibular functions of noise exposed rats were assessed by swimming tests. The morphology of vestibular hair cells was examined by scanning electron microscopy.

Conclusion
Conclusion: Our preliminary results indicated that vestibular deficits were induced by the unilateral noise exposure. Ongoing studies are to characterize the effects of air-conducted and bone-conducted noise exposure on the vestibular system and examine the underlying neural mechanisms.(supported by NIH R01 DC012060-01)
sensitivities and coherence measures in response to stimulus frequencies $\geq 1.6$Hz, these data suggest lower signal:noise ratios among responding afferents. The phase measures of some afferents' responses suggested they represented remnant dimorphic afferents projecting from the crista central zone. No evidence of induced lesions were observed subsequent to 0.5µg gentamicin administration, where the cristae and utriculi exhibited normal distributions of anti-calretinin and anti-KCNQ4 immunoreactive calyces. Afferent discharge characteristics in these specimens could not be distinguished from untreated specimens. Further analyses to reveal hair cell densities and additional cellular markers are ongoing.

Conclusion
These results demonstrate that partial lesions of the vestibular epithelia may be induced by quantities of gentamicin as low as 0.75µg, though the dosage window for producing lesions compatible for detailed study is surprisingly narrow. Calyces projecting to the crista central zones are most severely affected, yet recorded responses indicate that central zone hair cells are functional.

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Effect of Low Level Laser on Vestibular System after Ototoxic Damage
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Background
Already applied in numerous areas, laser-induced phototherapy is widely applied as a noninvasive treatment promoting cell recovery and repair processes. It is especially notable that low-level laser therapy (LLLT) has been approved by the U.S. Food and Drug Administration for the treatment of several diseases, including carpal tunnel syndrome and alopecia. In the present study, the ability to recover vestibular function after gentamicin exposure was assessed using an 830 nm diode laser

Methods
Gentamycin (110mg/kg/day) was injected intravenously for three days in 17 rats. Eight rats were irradiated (830 nm diode laser, transtympanic) in the left ear at an energy output of 200mW for 30 min for 7 days in a row (L group) and 9 rats served as a control (C group). Gain and asymmetry of Sinusoidal Harmonic Acceleration (SHA, 0.02-0.64Hz) were measured before the gentamicin exposure and also after the 1st, 3rd and 7th irradiations. After the 7th treatment, hair cells were observed using a confocal laser scanning microscope. The lateral canal ampullae were sectioned in the axis vertical to the crista ampullaris and 1 representative section were obtained per 50 $\mu$m (6 samples/ampulla). The sections were stained with phalloidin and DAPI, and the number of hair cells was counted for quantitative comparison.

Results
After 7 days of LLLT, the gain of the left ear (LLLT ear) was significantly higher than that of the right ear (non-treated ear) in the L group. Also, the asymmetry was deviated to the right side (15-33%), reflecting that the vestibular function of the left ear has recovered while that of the right ear has not. On the contrary, the gain of the C group did not recover in both ears after 7 days. The asymmetry was symmetrical (<±10%), implying that bilateral vestibular weakness did not recover in neither of the ears after gentamin exposure. The number of hair cells was 145.5±25 in the left ear (LLLT ear), and 139±25 in the right ear (non-treated ear) of the L group. Although the number of hair cells was larger in the left ear when compared with that of the right ear, it did not reach a statistical significance.

Conclusion
Low level laser treatment may promote the recovery of vestibular functions after systemic gentamicin injection. But whether this effect is via improved hair cell survival is not clear and more research is needed.
Title: Novel Cytoplasmic Extensions Link Basal Portions of Type II Hair Cells With Other Elements in Adult Mammalian Vestibular Epithelia

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Background
Mammals have two types of vestibular hair cells – type I and II - that differ with respect to their morphology, physiology, and innervation. A recent study of adult mice (Golub et al., 2012, J. Neuroscience, in press) showed that all hair cells that are spontaneously regenerated in utricles are type II-like and have branched extensions of their cytoplasm that reach several cell lengths. In this study, we examined undamaged utricles from adult mice and gerbils for evidence for this cell type in normal epithelia.

Methods
Vestibular end organs from 2-45 week-old mice or adult gerbils were dissected, fixed, and immunolabeled as whole-mounts with antibodies to the hair cell antigens, myosin VIIa, myosin VI, and calretinin and/or with antibodies to neural proteins (neurofilament, βII tubulin) and pre- and post-synaptic proteins (Ctbp2, PSD95, SV2, and ChAT). End organs were analyzed using confocal microscopy. Three-dimensional reconstructions of hair cells were generated using Figi image segmentation.

Results
Image analysis revealed that many type II hair cells in adult mice and gerbils have one or more cytoplasmic processes that project basolaterally from the each cell. Hair cell processes have a variety of shapes and sizes, ranging from oval “blobs” to long branched processes that extend several cell lengths. In mice, hair cell processes have a high density of Ctbp2-positive pre-synaptic ribbon components. They are also closely associated with post-synaptic specializations and ChAT immunolabeling, suggesting they receive inputs from efferent fibers. Three-dimensional reconstructions suggest that nearby type II hair cells seem to contact one another by their processes, and they confirm that type II hair cell processes contact calyces of type I hair cells. In mice, hair cell processes are first seen during the second postnatal week.

Conclusion
Many type II hair cells in mature rodents have a dramatic feature that has not been described before: one or more cytoplasmic processes that extend underneath type I hair cells, synapse on other cell types (most likely afferent nerves or hair cells), and receive synaptic input from other cells (most likely efferent nerves). Prior studies may have failed to notice this specific morphology of type II hair cells because they analyzed thin sections, young animals, or focused on type I hair cells. These observations suggest that direct communication amongst type II hair cells and cellular elements other than afferent fibers could modulate neural signaling in the vestibular system.
Persistent and Quantal Synaptic Currents in Vestibular Calyx Terminal

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Background

The unique morphological structure of the vestibular calyx has generated speculation concerning communication between Type I hair cells and the calyx inner leaflet membrane. Quantal and non-quantal transmission has been suggested. To examine this we recorded from calyces in the isolated lagenar macula of the turtle, Trachemys scripta elegans.

Methods

The lagenar epithelium was dissected, otoconia enzymatically removed and the tissue placed upon the fixed stage of an upright microscope for DIC and fluorescence imaging. Recording was performed employing the whole cell patch clamp technique. Patch electrodes were loaded with a fluorescent dye for viewing following recording. A fluid jet was employed to deflect hair bundles. The composition of the fluid in the jet was varied between a sodium-dominated perilymph like solution to a potassium-dominated endolymph like solution.

Results

All recordings were confirmed to have been taken from calyceal terminals by visualizing the terminal and its axon leaving the tissue employing wide field fluorescence imaging. Two types of excitatory post-synaptic currents (EPSC) were observed in voltage clamped vestibular calyx terminals held at -60mV. Miniature (mEPSCs) whose frequency could be increased by deflecting the hair bundle and an amplitude graded inward post-synaptic current that persisted for as long as the bundle was held deflected (pEPSC). pEPSCs developed and decayed with a time constant of ~ 200msec. Both types of EPSCs occurred simultaneously in a subset of cells. pEPSC varied from 0 to 200pA depending upon the cell and the intensity of bundle deflection. The frequency of mEPSCs adapted during prolonged bundle deflection while pEPSCs did not. Varying the solution in the fluid jet (cf. methods) did not affect the character of the responses. CNQX reversibly blocked the mEPSCs without affecting the pEPSC, while TTX reversibly blocked the pEPSC without affecting the mEPSCs.

Conclusion

Results imply that mEPSCs convey timing and high frequency information while pEPSCs convey low frequency and tonic signals (e.g., gravity), presumably at a much lower energetic cost to the hair cell. The fact that the pEPSC did not adapt to maintained bundle deflections suggests it reflects the tonic component of the transduction current.

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Sodium channel distribution in vestibular afferents

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Background

The sodium channel isoform Naᵥ1.6 is the canonical sodium channel at the node of Ranvier and the axon initial segment. We have been investigating Na channel distribution with various Pan-Naᵥ and Naᵥ isoform-specific antibodies in vestibular afferents to determine whether Naᵥ1.6 or some other isoform is the main isoform present at the AIS region of vestibular afferents.

Methods

Methods used were similar to those used in our recent publication (Lysakowski et al., 2011), with the exception that most tissue was vibratome-sectioned rather than frozen-sectioned and lower concentrations (0.05%) of Triton X-100 were used.

Results

Our results indicate that Naᵥ1.6 is present at the heminode of dimorphic vestibular afferents, but not at pure calyx afferent heminodes. We determined the afferent class and location of the heminode by co-labeling with calretinin (a calyx afferent marker) and nodal marker antibodies, such as ezrin, βIV-spectrin, neurofascin-186 and ankyrinG, and paranodal and juxtaparanodal marker antibodies, such as Caspr1, ankyrinB, Kᵥ1.1 and Kᵥ1.2, and myelin basic protein. Our search for the Naᵥ isoform at calyx heminodes led us to investigate various other isoforms, including Naᵥ1.1-1.3, 1.5, 1.7 and 1.8, in addition to various Pan-Naᵥ antibodies. Each antibody labels subsets of heminodes, although we have not yet quantified the proportions of individual isoforms expressed within each afferent class. Pan-Naᵥ and Naᵥ1.2 label the entire lower
outer surface of the calyx in dimorphic afferents, down to the heminode. Calyx afferents are labeled with Na\textsubscript{V}1.3 and Na\textsubscript{V}1.8 antibodies in a similar pattern. The majority of calyx afferent heminodes are located above and the majority of dimorphic afferent heminodes below the basement membrane.

Conclusion
We conclude that while Na\textsubscript{V}1.6 has a role as the major player at the vestibular dimorphic afferent heminodes, other isoforms have supporting roles.

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A Constitutive Null Mutation for Calretinin Has No Effect on Excitability of Vestibular Ganglion Neurons
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Background
Work with transgenic animals has shown that calretinin, a fast calcium binding protein, can modulate the electrical activity of certain CNS neurons. Calyx-only vestibular ganglion neurons (VGNs), which innervate type I cells in central zones of the sensory epithelia, express high levels of calretinin. The calretinin-positive, calyx-only afferents have irregular firing, in contrast to the regular firing of calretinin-negative afferents to peripheral zones. We used calretinin-null mice (obtained from S. Schurmans) to investigate whether calretinin plays a role in the firing properties of calretinin-positive calyx-only VGNs.

Methods
VGNs were isolated from mice (60-90 days old) lacking functional calretinin (CR -/-), heterozygotes (CR +/-), and their wild-type littermates (CR +/+), and cultured overnight. Whole-cell currents and voltages were recorded with the perforated patch clamp method.

Results
Two groups of VGNs were identified based on their responses to depolarizing current steps between 50 and 500 pA: “transient” spikers made one-two spikes at step onset and “sustained” spikers made more than two spikes. As reported for VGNs from young rats (P0-P14), transient spikers are larger and have more negative resting potentials and larger current thresholds for spiking. Previous work associated the transient firing pattern with irregular afferents, which include the calyx-only, calretinin-expressing afferents, and the sustained firing pattern with regular afferents, which are calretinin-negative (Kalluri et al., J Neurophysiol. 104:2034, 2010). We hypothesized that effects of the calretinin null mutation would manifest most strongly in transient spikers. Neither transient nor sustained spikers, however, showed significant differences with genotype in spike parameters (height, width, current and voltage thresholds, afterhyperpolarizations) or firing patterns.

We also attempted to calibrate the endogenous Ca\textsuperscript{2+} buffering of VGNs, using spike properties and firing patterns as assays. After recording in perforated-patch mode to determine firing properties in endogenous buffering, we ruptured the membrane and replaced endogenous buffering with the fast synthetic Ca\textsuperscript{2+} buffer, BAPTA. For all 4 BAPTA concentrations used (0.002, 0.1, 1, and 10 mM), membrane rupture was followed by a small depolarization and small effects on spike properties and firing pattern. Because these effects did not vary with BAPTA concentration, we expect that they reflect some other difference between the endogenous intracellular solution and the pipette solution.

Conclusion
VGN spike properties were resistant to constitutive changes in calretinin expression and acute changes in Ca\textsuperscript{2+} buffering.

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Distribution of 5-HT1F receptors in monkey vestibular ganglia
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Background
Migraine and balance disorders are often comorbid with psychiatric disorders. Clinical data suggest that serotonergic mechanisms play an important role in vestibular contributions to migraine and balance disorders. Both 5-HT1B and 5-
HT1D receptors are expressed extensively in inner ear ganglion cells of monkeys and rats. This study describes the distribution of another target of triptans, the 5-HT1F receptor, in vestibular ganglion cells of monkeys.

**Methods**
Decalcified, paraffin embedded temporal bone sections (8 – 10 microns) from 5 adult macaques monkeys were stained immunohistochemically for 5-HT1F receptors with a diaminobenzidine chromagen. Polyclonal, primary antibodies (Imgenex 78120/78121, 1:1000-5000) recognized human 5-HT1F N- or C-terminus. A series of 12 bit digital images were taken with a 20X objective and neuronal somatic area and intensity of immunoreactive vestibular ganglia were quantified using Meta-Morph software. Data were normalized by subtracting background values and analyzed statistically employing SYSTAT 11 (SYSTATSoftware, Inc.).

**Results**
Virtually all vestibular ganglion cells were immunopositive for 5-HT1F receptors, with staining confined to distinct cell regions. Staining of inferior and superior vestibular ganglia was more intense in small cells, punctuate in some medium cells and polarized regionally in some large cells. Analyses of average somatic neuronal immunoreactive intensity identified a population of mainly medium sized cells with high standard deviation of intensity corresponding to punctately stained cells. Centrally directed ganglion cell processes were immunoreactive as well as ganglion cell dendrites, cochlear inner hair cells, vestibular hair cells and blood vessels in vestibular maculae and cristae.

**Conclusion**
Although the 5-HT1F, 5-HT1B and 5-HT1D subtypes are expressed in almost all vestibular ganglia, they show slightly different cellular distribution patterns. Interestingly, 5-HT1F receptors are expressed in analogous regions in trigeminal, dorsal root and vestibular systems. We therefore propose that 5-HT1F receptors may act in concert with other subtypes in vestibular ganglia, in mediating serotonergic involvement in the comorbidity of migraine and balance disorders. In addition, 5-HT1F receptor expression in cochlear inner hair cell afferents suggests that this receptor could be exploited as a pharmacological target in treatment of audiovestibular symptoms of migraine.
Are There Differences in Infectability, Reactivation, or Production of Herpes Simplex Type I Virus in Superior versus Inferior Vestibular Ganglion Neurons?

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Background
Vestibular neuritis is an extremely common condition causing spinning vertigo, which results in at least 15% of all complaints of dizziness to primary care physicians. In these patients, involvement of the inferior vestibular ganglion is rare (36% or less), and exclusive involvement is even less common (5%). Many authors have found evidence that vestibular neuritis results from herpes simplex type I (HSV1) infection or reactivation in the vestibular ganglion. In this study, we used an in vitro cell culture system of herpes infection and reactivation to determine whether there are intrinsic differences in HSV1 infection, reactivation, or virion production in inferior versus superior vestibular ganglion neurons (VGNs).

Methods
Primary cultures of rat superior and inferior vestibular ganglion neurons were cultivated separately. Neurons were lytically and latently infected with HSV1 modified with a US11-green fluorescent protein (GFP) chimera. Percent lytic infection and baseline reactivation rates were assessed by fluorescent microscopy for GFP fluorescence. Trichostatin A (TSA) was applied to cultures and reactivation of HSV1 was assessed by fluorescent microscopy. Production of HSV1 virions was assessed by viral titer on Vero cells. The number of cells latently infected for both types of neurons was determined by fluorescent in situ hybridization for LAT transcripts.

Results
Lytic infection rates were equivalent between the two ganglia, with 100% lytic infection rate at an multiplicity of infection of 3. Lytic infections yielded similar amounts of plaque forming units (pfu; 3.25 x 10⁵ v. 6.6 x 10⁵ pfu/mL). Numbers of LAT positive neurons by FISH were not significantly different for superior and inferior ganglia. Latently infected superior and inferior ganglion cultures showed no differences in rates of baseline (16.8% v. 17.9%, p>0.05) and TSA-induced (64.3 v. 59.7%, p>0.05) HSV1 reactivation as measured by GFP fluorescence. Production of HSV1 virions was also not different between reactivated, latently infected superior v. inferior VGNs (3.5 x 10⁴ v 2.6 x 10⁴ pfu/mL).

Conclusion
Differences in the prevalence of superior and inferior vestibular neuritis are unlikely to result from intrinsic differences in HSV1 infection or virion production between the neurons contained in these ganglia. Other factors, such as the length and width of the bony canal containing the ganglia and nerves and accessibility of the ganglia by cerebrospinal fluid, may account for the discrepancy in involvement of the two ganglia.
Functional analysis of TRPV1 and TRPA1 channels in rat vestibular ganglia

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Background
Transient receptor potential vanilloid (TRPV)-1 and transient receptor potential ankyrin (TRPA)-1 are both cation channels co-expressed in sensory neurons such as dorsal root ganglia (DRG) and trigeminal ganglia (TG). TRPV1 is activated by noxious heat (>43°C), protons, capsaicin, and anandamide to name a few, while TRPA1 is activated by noxious cold stimuli (<17°C), alkaline pH (>8.5), cinnamaldehyde, mustard oil, allyl isothiocyanate and icilin to name a few. TRPV1 has been reported to be expressed in vestibular ganglia (VG) and suggested to be associated with vestibular function and/or dysfunction. On the other hand, in DRG and TG neurons, TRPV1 is co-expressed with TRPA1, but expression of TRPA1 in VG neurons has not been reported yet. Moreover, it is unclear whether TRPV1 and TRPA1 are functional as ion channels.

We studied the histological and functional expression of TRPV1 and TRPA1 in rat VG neurons.

Methods
We examined the TRPV1 and TRPA1 mRNA expression and localization by RT-PCR and in situ hybridization experiments and their functional analysis by Ca2+-imaging experiments.

Results
RT-PCR specifically amplified TRPV1 and TRPA1 transcripts from rat VG. In situ hybridization experiments showed TRPV1 expression in majority of VG neurons and also TRPA1 expression. In Ca2+-imaging experiments, capsaicin, a TRPV1 agonist, induced significant increases in intracellular calcium ion concentration ([Ca2+]i) in rat primary cultured VG neurons, which were almost completely blocked by capsazeine, a TRPV1 selective antagonist. Cinnamaldehyde, a TRPA1 agonist, also caused [Ca2+]i increase, which were completely inhibited by HC030031, a TRPA1 specific antagonist. Moreover, in some VG neurons, [Ca2+]i increase was evoked by both capsaicin and cinnamaldehyde in the same neurons. We examined a total of 114 VG neurons and 21.9% (25/114) responded to capsaicin, 28.1% (32/114) to cinnamaldehyde, and 12.3% (14/114) both to capsaicin and cinnamaldehyde.

Conclusion
In summary, our data showed histological and functional expression of TRPV1 and TRPA1 in VG neurons. Ca2+-imaging experiment suggested that these two channels can be co-expressed in the same neuron. It is suggested that TRPV1 and TRPA1 in VG neurons might participate in vestibular function and/or dysfunction such as vertigo.
Opioid receptor mediated inhibition of the calcium currents in vestibular ganglion neurons of the rat

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Background
Opioid receptors have been shown to be expressed in vestibular endorgans, but studies of their mechanism of action at cellular level are scarce. With the aim to determinate the opioid receptors function we evaluated the modulation of the voltage-gated calcium current ($I_{Ca}$) by the $\mu$ opioid receptor activation (MOR) in afferent neurons isolated from the vestibular ganglia of the rat.

Methods
Young (P7-10) and prepubertal (P28-30) rats were studied using the perforated whole cell voltage clamp technique.

Results
The use of MOR agonists [D-Ala², N-Me-Phe⁴, Gly⁵ -ol]-enkephalin (DAMGO), Met-enkephalin or endomorphine-1 inhibited the $I_{Ca}$ in vestibular neurons at both ages studied. The DAMGO inhibition was dose-dependent with an IC₅₀ of 33 nM. The action of DAMGO was prevented by the co-application of MOR antagonist H-D-Cys-Tyr-D-Trp-Arg-Thr-Pen-Thr-NH₂ (CTAP) or in neurons pre-incubated with pertussis toxin (18-24 h, 500 ng/ml). Co-application of 8-Br-cAMP with DAMGO, reverted the inhibition induced by DAMGO and recovered 92 % of the control current, indicating that cAMP is the second messenger involved in the inhibition of $I_{Ca}$ by MOR. The application of the protein kinase A inhibitor H-89 mimicked the DAMGO effect, and the co-application of H-89 and DAMGO did not add their effects. In contrast, the co-application of porbol-myristate was unable to recover the DAMGO inhibition. DAMGO inhibited the LVA and HVA currents. The use of nickel, nifedipine and $\omega$-ctx-GVIA showed that DAMGO inhibited both the low and the high voltage activated Ca²⁺ currents. In current clamp experiments the MOR activation increased the duration and decreased the amplitude of the action potential and decreased the repetitive discharge in response to depolarizing current pulse injection.

Conclusion
We concluded that in isolated vestibular afferent ganglion neurons the $\mu$ opioid receptor inhibits the T-, L- and N- calcium current, trough the activation of a Gᵢ/o protein that leads to a decrease of the cAMP levels and the subsequent decrease of the PKA activity. The negative modulation of the Ca²⁺ current would modify the postsynaptic signal integration in the synapse between the hair cell and the vestibular afferent neuron and also at the central level decreasing the neurotransmitter release in the synapse between the vestibular afferent neurons and vestibular nucleus neurons.
Modification of the Hodgkin-Huxley Equations According to Experimental Data from Vestibular Afferent Neurons.
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Background
Development of the vestibulo-sensorial conflict in microgravity leads to increased delays of the gaze fixation reaction (3 fold time increase in comparison with terrestrial conditions). The use of a micro accelerometer and a micro gyroscope, established on a helmet of a cosmonaut, are thought to help reduce such delay development. The use of a processing unit of the information from technical devices will provide the adequate correcting signals.

Methods
In this work the afferent neuron discharge is modeled on the basis of a Hodgkin-Huxley (HH) modified differential equations system.

The functional parameters and numerical data used in the model have been obtained from experimental measurements in the vestibular afferent neurons of the rat.

Results
The novelties of the simplified system that we are proposing are the following:
1. In the description of the potassium current ($I_k$) we have introduced a parameter of inactivation $h_k$ of the $I_k$ and in this connection we use one more equations of Kolmogorov for the description on the average dynamics of this parameter.
2. It has been mathematically justified the reduction of an order of the mathematical model in the presence of a small parameter $\delta m$ at a derivative of the subsystem describing on the average dynamics of activation parameter $m$ of $I_k$.
3. The integral $n + h = c = \text{const}$, where $n$ is the activation parameter of the $I_k$ and $h$ is the inactivation parameter of the sodium current ($I_{Na}$), were used as a simplification of the HH equations in various models (Rubin, 2000). In our work it is modified according to experimental data as $n + h = c (v)$ where $v$ is the voltage.

Conclusion
Thus, modified model is presented in the form of Cauchy equations of the third order with small parameter in the right part of last equation describing on the average dynamics of the new parameter $h_k$.

The mathematical analysis of the model leads to conclude that this can be used as a part of a complete model of information processing in the vestibular end organs, that can be numerically solved using a microprocessor and used for the design of an automatic correction of the gaze stabilization in extreme conditions such as microgravity.

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Afferent Responses to Efferent Stimulation in the Murine Vestibular System
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Background
The vestibular organs of nearly every vertebrate receive a prominent efferent innervation whose activation modulates the ongoing discharge of vestibular afferents. Responses of mammalian vestibular afferents to efferent stimulation are effectively excitatory and have been extensively characterized in squirrel monkey and chinchilla. However, there is little data regarding the underlying cellular and synaptic mechanisms. The use of transgenic mouse models in tandem with pharmacological manipulation provide a powerful opportunity to parse out the various receptors and downstream effectors involved in mammalian efferent responses. The purpose of this study was to begin characterizing the response of mouse vestibular afferents to stimulation of efferent neurons.

Methods
C57Bl6 mice were anesthetized with urethane/xylazine, fitted with a tracheal cannula, and connected to a ventilator. A posterior craniotomy and cerebellar aspiration were performed to expose the right vestibular nerve and floor of the fourth ventricle. Glass microelectrodes were used to record the discharge of vestibular afferents in CNVIII, visually identified just as it exits the otic capsule. The discharge regularity (CV*) of each afferent was computed from interspike intervals taken from background activity. To electrically stimulate vestibular efferents, shock trains (1-5s, 200-333/s) were delivered to an electrode array consisting of four wires spaced 400 µm apart. The array was lowered into the floor of the fourth ventricle along the midline. The facial nerve was transected bilaterally and/or tubocurarine was administered IP to block muscle contractions.

Results
We have recorded from a total of 26 units. Similar to that reported in squirrel monkey and chinchilla, mouse vestibular afferents were excited following midline efferent stimulation. This excitation consisted of a modest, fast perstimulus excitation (10-30 spikes/s) and slower excitation persisting for 50-60 seconds after the stimulus. Efferent responses were routinely observed in regularly to moderately regularly-discharging afferents (CV*<0.15) whereas most of the responses in irregularly-discharging afferents (CV*>0.15) were unexpectedly small or absent. These observations may be related to the stimulus location, as midline placement of the array would be predicted to stimulate predominantly contralateral efferent fibers, presumed from studies in other species to innervate mostly regular afferents.

Conclusion
Like previous mammalian studies, mouse vestibular afferents are excited by the midline activation of efferent neurons. The resulting excitation consists of fast and slow components differing in both their activation kinetics and duration. These results will be compared to those obtained with the additional stimulation of ipsilateral efferent neurons.

Loss of alpha-Calcitonin Gene-Related Peptide (CGRP) reduces Vestibuloocular Reflex (VOR) gain
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Background
All hair cell systems contain an efferent innervation originating in the brainstem and projecting to the hair cells and/or afferent processes innervating the sensory epithelium. However, the functional role of the brain's efferent projection back to the vestibular component of the inner ear remains a mystery. Previous studies have characterized the neuroanatomy of this efferent vestibular system (EVS) projection, yet have not revealed its role in modulating vestibular system function. Calcitonin-gene related peptide (CGRP) is a neuroactive peptide known to act at the synapses between vestibular and cochlear efferent fibers and their targets. In the cochlea, we and others have found that lateral olivocochlear fibers contain CGRP and have determined that the loss of CGRP in knock-out mice reduces sound-evoked activity of the cochlear nerve by ~20%. Because CGRP is also an EVS neuropeptide, we have begun to investigate whether and how the loss of CGRP influences vestibular system function in CGRP null (−/−) mice.

Methods
We determined that in CGRP null (−/−) mice, CGRP staining was absent in all vestibular canals and extraocular muscles using immunohistochemistry to detect CGRP. In contrast, AChE staining in the canal and extraocular muscles was comparable in normal and CGRP null (−/−) mice. To test vestibular function, mice were secured with the horizontal semicircular canals level with the horizontal plane and eye movements were recorded using an infrared camera system.
Turntable velocity was measured and data acquisition was controlled online. To elicit the VOR, mice were rotated in the yaw axis at frequencies of 0.2-4 Hz, at a peak velocity of 15º/s.

Results
We have found that the loss of CGRP drastically reduced the VOR gain by ~50%. However, one alternative explanation for the deficit observed in CGRP null (−/−) mice is that it reflects a non-vestibularly related effect of the CGRP removal at the level of the extraocular motor neurons and/or their innervations of the extraocular muscles. We have tested this possibility indirectly using immunohistochemistry, and directly by confirming that the relationship between vestibular quick phase amplitude and its peak velocity is comparable in normal and CGRP null (−/−) animals.

Conclusion
The efferent vestibular system (EVS) neurotransmitter CGRP plays a key role in the development of the VOR gain. Information from these studies may also contribute to future drug therapies aimed at enhancing vestibular function.

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Sensitivity of semicircular canal afferents to temperature gradients and pulsed infrared radiation

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Background
In the present study we aimed to determine the effect of changes in temperature induced at the vestibular neuroepithelium on the resting discharge rate and sensitivity of semicircular canal afferents. The temperature gradients were introduced by perfusion of temperature controlled artificial perilymph and with pulsed infrared radiation. In addition we have also examined the role of inhibitory and excitatory convergent synaptic inputs from the hair cells to the exquisite temperature sensitivity of the vestibular labyrinth.

Methods
The experiments were conducted in the oyster toadfish, Opsanus tau and responses of single afferent fibers were recorded. The temperature changes were generated by perfusion of the entire labyrinth with temperature-controlled artificial perilymph solution. The temperatures were controlled between 5º to 38º. In addition, the responses of the afferents were recorded in response to pulsed laser radiation (Capella laser, Lockheed Martin Aculight, λ = 1860nm). The fish core temperature was maintained at ambient using a sea-water bath. The changes in spontaneous discharge rate, their regularity, adaptation to step hair bundle displacements and responses to sinusoidal head rotations of single-units were recorded. The roles of GABAergic and Glutamatergic hair cell inputs to the temperature-controlled responses were examined. Following fixation, epithelia containing injected afferents were processed for two-color fluorescence for biocytin and GABA, and whole mounts were imaged by two-photon microscopy.

Results
A variety of afferent responses to thermal perfusion and infrared irradiation were observed. Some of the recorded afferents showed an increase in the resting discharge rate with increased temperature while the others reduced their discharge rate. The responses to pulsed infrared radiation varied in the same afferents with increased and reduced temperatures. The arborization patterns of the vestibular afferents receiving GABARergic and Glutamatergic hair cells inputs were studied.

Conclusion
The results indicate that the convergent hair cell inputs may play a role in the temperature sensitivity of the vestibular labyrinth.

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Vestibulo-Ocular Reflex Eye Movement Responses to Infra-Red Laser Stimulation of the Mammalian Labyrinth

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Background
The aim of this study was to determine whether IR stimulation can evoke vestibulo-ocular reflex (VOR) responses in mammals. Rajguru et al. recently reported that pulsed infrared (IR) laser stimulation of hair cells can modulate afferent fiber activity in neurons innervating the crista ampullaris of the toadfish vestibular labyrinth. Extending this approach to mammals, Ahn et al. demonstrated similar afferent responses to IR stimulation of the crista of chinchillas (C. lanigera). Since these single-unit electrophysiologic responses included a mix of large excitatory, inhibitory, and mixed responses from canal afferents apparently independent of caloric-induced endolymph convection, whether prosthetic IR stimulation can evoke significant VOR eye movement responses has been unclear.

Methods
Using the same approach to fiber placement when measuring single-unit electrophysiologic responses of superior and horizontal canal afferents during IR stimulation, the left bullae of adult, wild-type 450-650 g chinchillas were opened under isoflurane general and local field block anesthesia. A hole was drilled in the superior canal thin segment at its junction with the ampulla. A 400 µm optical fiber was positioned in the superior canal and oriented inferomedially (toward the superior and horizontal canal cristae). Isoflurane was titrated down until VOR responses to whole-body rotation about the horizontal canal axis in darkness were recovered. IR stimulation was then delivered by a Capella Infrared Neural Stimulator (Lockheed-Martin Aculight, Bothell, WA, USA) at 1870 nm wavelength, with repetition rates of 200-400 pulses/s, and pulse widths of 200–350 µs.

Results
In two out of four chinchillas, eye movements were observed during IR stimulation of the labyrinth, with peak slow phase velocity exceeding 15°/s and direction consistent with left horizontal semicircular canal inhibition.

Conclusion
Stimulation of the mammalian vestibular labyrinth using infrared laser stimulation produced VOR responses consistent with modulation of activity in the horizontal canal. Whether this is due to cupular deflection (e.g., via convection) or a more direct action on hair cells and/or afferents is not yet known, but can be determined by characterizing the dependence of responses on head orientation and on the status of hair cells after ototoxic injury. Further study of IR as an alternative means of prosthetically stimulating the mammalian vestibular labyrinth is warranted.

Supported by a research contract from Lockheed-Martin-Aculight
Introduction to the Workshop - "Assessing Tinnitus in Animal Models: Progress and Pitfalls"
M. Charles Liberman

Understanding tinnitus mechanisms requires work in animal models; however, the measurement of a phantom percept in an animal presents obvious challenges. Several techniques for tinnitus assessment have been developed, including conditioned avoidance to foot shock, and gap-prepulse inhibition of the acoustic startle reflex. This Workshop aims to improve understanding of the strengths and weaknesses of the different approaches, and to promote the refinement of their application.

1) Two 20-minute talks will introduce the topic. The first will summarize the psychophysics of tinnitus in human subjects. The second will review the behavioral strategies devised to assess the presence of tinnitus in animals, as well as the evidence for which parts of the brain each test engages, and the existing data on cross-test validation.

2) Next, two 15-minute talks will describe specific applications of the tests: a) one conditioned avoidance test and one startle-reflex test. Each speaker, who is also the test’s developer, will describe test design, rationale, training requirements and reproducibility of results.

3) Lastly, a moderated 60-minute panel discussion will include the speakers, a representative of the Tinnitus Research Consortium (http://www.ncbi.nlm.nih.gov/pubmed/22245715) and other tinnitus researchers. Audience members can bring 1-2 relevant slides to frame their question and/or facilitate discussion, with a time limit of 3 minutes per participant.

Behavioral Strategies for Tinnitus Assessment in Animals
Brad May

Animal models provide a critical context for controlled laboratory studies of the physiological mechanisms of tinnitus. A longstanding challenge for this research is verification of phantom sound perception in non-human subjects that cannot verbally report the subjective properties of experimentally induced tinnitus. This presentation will introduce the types of behavioral strategies that have been used for tinnitus assessment in animals. We will begin by summarizing the basic experimental methods for inducing tinnitus. Then, we will demonstrate how the altered perception of silence can serve as an objective behavioral marker of tinnitus status. We will describe how the context of behavioral testing can be arranged to quantify additional subjective characteristics of tinnitus, such as its pitch and magnitude. The validity of these animal models will be assessed by comparing outcomes that have been obtained with different behavioral approaches from animals that have been treated with comparable induction methods. As tinnitus investigators seek high-throughput behavioral methods for screening animals, reflex-based methods have supplanted conditioning paradigms that may demand a substantial time investment in each subject. A concern raised by these new approaches is whether a change in reflexive behavior adequately reflects abnormal auditory processing in the parts of the brain that relate most directly to tinnitus perception.

Psychophysics of Tinnitus in Human Subjects
James Henry

Over 30 years ago formal efforts were undertaken by the CIBA Foundation in London to promote international cooperation in tinnitus research. A central concept in the CIBA Symposium was that standardization of tinnitus measures would advance international understanding and facilitate work on tinnitus. As a result of these efforts a clinical assessment battery was recommended to include pitch match, loudness match, maskability, and residual inhibition. Many tinnitus researchers and clinicians routinely obtain these measures. However, normative data do not exist to facilitate interpretation of the measures.

Methods
We have conducted studies since 1995 to develop computer-automated, self-guided tinnitus assessment procedures that can be used in a standardized fashion. The automated system has undergone numerous iterations and was eventually redesigned to enable clinical implementation. The Tinnitus Evaluation System (TES) now comprises a laptop computer
Results
Measures obtained with the TES included (1) hearing threshold; (2) loudness match; (3) pitch match; (4) bandwidth match; (5) minimum masking level; and (6) residual inhibition. Repeated testing with research participants has documented response reliability for all of these tests. In general, all of these measures are reliable upon repeated testing except for pitch matching.

Conclusion
Normative data are important to identify relationships between measures of tinnitus perception and other factors that could help to: (1) elucidate underlying mechanisms of tinnitus; (2) facilitate a more definitive assessment of the association between these measures and measures of tinnitus impact; and (3) predict specific types of therapy that might be most beneficial to an individual patient.

A Startle-Reflex Based Test for Tinnitus
Jeremy Turner

The startle reflex, first explored systematically in the late 1930s by Landis and Hunt, has developed into a valuable tool for behavioral neuroscience research with many applications. The reflex and its habituation are routinely used to assess sensorimotor adverse effects in preclinical toxicology testing and drug development. The reflex can also be elevated through fear conditioning (known as fear-potentiated startle) and this technique has been used for several decades in the search for the neural basis of fear and anxiety, as well as in the development of anxiolytics. Another valuable feature of the startle reflex is how preceding sensory cues presented before the startle stimulus (optimally 50-150 ms), have the ability to reduce, or inhibit the reflex. This feature, generally referred to as prepulse inhibition (PPI) of the startle reflex, has been used to conduct audiometric/psychophysical testing in animal models (most notably by the laboratories of Ison, Willott, and their colleagues). PPI has also been used to document sensory processing deficits in human and animal models of schizophrenia and related disorders. A recent adaptation of the startle reflex test has been proposed for the measurement of tinnitus. In this test, the startle reflex is inhibited when a silent gap embedded in a background noise is used as an inhibitory preceding cue. The hypothesis is that if an animal has ringing in the ears the silent gap cue would serve as a less salient stimulus, and thus a less effective inhibitor of the subsequent startle reflex. This presentation will review the use of the gap technique for measuring tinnitus and will discuss what has been learned about its applications. We will also discuss some of the challenges in using this technique and some of the questions that still remain.

A Conditioned-Avoidance Based Test for Tinnitus
Shaowen Bao, Sungchil Yang, Benjamin Weiner, Li Zhang

Tinnitus is conscious perception of phantom sounds. We developed a behavioral test for tinnitus in a rodent model, in which this conscious perception drives a volitional active avoidance response (as compared to an autonomous, reflexive response). Our rationale is based on the assumption that while reflexes could be elicited subconsciously, volitional responses require decision-making that is based on conscious sensory perception. The behavioral procedure is divided into a training phase and a testing phase. In the training phase, animals learn to avoid foot shocks by crossing a barrier that separates the two compartments of a shuttle box when any sound is played. After training, animals are subjected to noise-induced hearing loss (NIHL) and begin the testing phase. A post-NIHL increase in barrier-crossing behavior in the absence of a sound is taken to indicate the presence of a tinnitus percept. This tinnitus measure is reproducible. Furthermore, it is correlated with the reduction of cortical GAD65 expression, supporting a role of the auditory cortex in the presumptively conscious tinnitus perception.
Optogenetics: Molecules and Hardware for Control of Neurons
Edward Boyden

We have developed tools for the optical control of specific cell types embedded within the nervous system. These tools are now being used scientifically to reveal new targets for treating neural disorders, and also may clinically support prototype treatments for diseases ranging from blindness to post-traumatic stress disorder. A key aspect of our prosthetics is our development of a suite of fully-genetically-encoded reagents (e.g., ChR2, Halo, Arch, Mac, ArchT, Jaws, Chronos, and others) that, when expressed in specific neuron types in the nervous system, enable their activities to be powerfully and precisely activated and silenced in response to pulses of light. These tools are in widespread use for analyzing the causal role of defined cell types in normal and pathological brain functions. In this talk I will briefly give an overview of the field, and then I will discuss a number of new tools for neural activation and silencing that we are developing, including new molecules with augmented amplitudes, novel color and light-sensitivity capabilities, and unique new capabilities. I will also describe new devices for high-channel-count light delivery into the awake behaving nervous system. Finally, I will talk about how these tools can be used to enable systematic analysis of neural circuit functions in the fields of emotion, sensation, and movement, and in neurological and psychiatric disorders, as well as our pre-clinical work on translation of such tools to support novel ultraprecise neuromodulation therapies for brain disorders.

Luminopsin and Clomeleon: Novel Optogenetic Tools for Controlling and Monitoring Neuronal Activity
Ken Berglund

Over the past several years optical tools for observing and controlling neuronal activity have vastly expanded, from genetically encoded optical reporters to various light-driven actuators. However, there are unmet needs which require developing novel tools. We have developed two such tools, namely luminopsin and Clomeleon. I will provide proofs of principles for the two techniques and summarize their applications in neuroscience research.

First, it would be advantageous to manipulate neuronal activity acutely and invasively as well as chronically and non-invasively, using the same genetic construct. Luminopsin allows activation of genetically defined populations of neurons by means of localized physical light and a systemically applied chemical using the same construct. Luminopsin was created by fusing a light-sensing channelrhodopsin with a light-emitting protein, Gaussia luciferase. The channelrhodopsin in this chimeric molecule can be activated by physical light (standard optogenetic approach) as well as by light generated by the luciferase acting on its substrate coelenterazine, CTZ (chemical genetic approach). We transiently transfected cultured neurons with luminopsin constructs and demonstrated the feasibility of using CTZ-activated bioluminescence to modulate ongoing neuronal activity. Specifically, application of CTZ depolarized the membrane potential of luminopsin-expressing neurons, which enhanced neuronal excitability and caused neurons to fire more action potentials in response to synaptic input. Thus, luminopsin expands the current approaches of manipulating neural activity in the brain by allowing channelrhodopsins to be activated by a diffusible molecule as well.

Second, while there are several approaches to image synaptic excitation and action potential firing in neurons, synaptic inhibition has remained largely invisible, partly due to a lack of appropriate indicators. Clomeleon is an optogenetic tool that was intended to shed a light literally to synaptic inhibition mediated by chloride (Cl). Clomeleon is a fusion protein of Cl-sensitive yellow and Cl-insensitive cyan fluorescent proteins. Through fluorescence resonance energy transfer (FRET), it enables ratiometric measurements of intracellular Cl concentration, which reflects and dictates synaptic transmission mediated by Cl. We developed and characterized transgenic mouse lines that express Clomeleon in different parts of the brain. Specifically, we have utilized these mice to demonstrate developmental Cl change in immature neurons and both phasic and tonic forms of synaptic inhibition in more mature neurons in the hippocampus and the cerebellum, respectively. In conclusion, Clomeleon provides an array of unique applications to address Cl regulations in neurons and possibly in other cell types in the least invasive manner.
High-speed Optogenetic Mapping of Brain Circuitry
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Although the basic anatomy and cellular components of the brain have been known for more than a century, we still do not understand how the brain works. This is in large part due to technical limitations in our ability to identify the function of brain circuits. New optogenetic technologies promise to greatly increase our understanding of brain circuit function.

By scanning small spots of laser light, synaptic circuits can be mapped in brain slices from transgenic mice expressing channelrhodopsin-2 (ChR2). These light spots photostimulate presynaptic neurons expressing ChR2, while postsynaptic responses can be monitored in neurons that do not express ChR2. Correlating the location of the light spot with the amplitude of the postsynaptic response elicited at that location yields maps of the spatial organization of the synaptic circuits. This approach yields maps within minutes, which is 10,000 times faster than can be achieved with conventional paired electrophysiological methods.

We have applied this high-speed technique to map local circuits in many brain regions. In cerebral cortex, we observed that maps of excitatory inputs to pyramidal cells were qualitatively different from those measured for interneurons within the same layers of the cortex (PNAS 104: 8143). In cerebellum, we could use this approach to quantify the convergence of molecular layer interneurons on to Purkinje cells and found that at least 7 interneurons form functional synapses with a single Purkinje cell. The number of converging interneurons is reduced by treatment with gap junction blockers, indicating that electrical synapses between interneurons contribute substantially to the spatial convergence. Remarkably, gap junction blockers affect convergence in sagittal cerebellar slices but not in coronal slices, indicating sagittal polarization of electrical coupling between interneurons. By measuring limb movement, this approach also can be used in vivo to define motor maps non-invasively (J. Neurosci. Meth. 179: 258).

In summary, ChR2-mediated high-speed mapping promises to revolutionize our understanding of brain circuitry.

Gating thalamocortical synaptic plasticity and auditory cortical map plasticity after the early critical period
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Gating thalamocortical synaptic plasticity and auditory cortical map plasticity after the early critical period
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The learning of behaviorally important sensory information is accompanied by an expansion of cortical representation areas in the sensory cortices. The substrate of this cortical plasticity is unknown. In the auditory cortex of neonates, cortical plasticity is induced by passive exposure to tones, but in adults, it requires active learning or pairing tones with the activation of cholinergic inputs. Like cortical plasticity, long-term potentiation (LTP) and long-term depression (LTD) at thalamocortical excitatory synapses can be induced in neonates but not in adults. Using single-cell electrophysiology, two-photon glutamate uncaging, and optogenetics in thalamocortical slices containing the auditory thalamus and auditory cortex, we determined that postsynaptically expressed LTP and LTD at thalamocortical synapses in the auditory cortex can be unmasked in mature mice by cortical disinhibition and activation of cholinergic projections emanating from the nucleus basalis. Cholinergic projections activate M\textsubscript{1} cholinergic receptors (M\textsubscript{1}Rs) at thalamic presynaptic terminals and sustain glutamate release from thalamic projections to the auditory cortex via negative regulation of adenosine signaling. Experiments in slices and in vivo showed that deletion of A\textsubscript{1} adenosine receptors bypasses the M\textsubscript{1}R-dependent requirement for LTP and LTD and permits cortical plasticity in the auditory cortex in adult mice, which is induced by passive tone exposure. These data suggest that synaptic plasticity at thalamocortical synapses underlies experience-dependent cortical plasticity in sensory cortices.
A comparison of electrical and optical activation of midbrain and cortical pathways in mice expressing channelrhodopsin-2 in the cochlear nucleus

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Background:

The majority of auditory brainstem implant (ABI) users have reduced performance compared to cochlear implant users. One possible explanation for this discrepancy is that existing ABI electrode designs are based on electrical stimulation and may have significant current spread and poor selectivity. This reduces the number of effective acoustic channels and increases side effects, such as dizziness or facial twitching from activation of non-auditory centers. Optogenetic activation of neurons (using channelrhodopsin-2, ChR2) has been used successfully in a number of systems as a way to improve selectivity of stimulation compared to electrical current. We hypothesize that cochlear nucleus (CN) neurons transfected with AAV-ChR2 can mediate responses along the ascending auditory pathways.

Methods:
The CN of CBA/J mice were virally-transfected with AAV2/ChR2. Pressure injection of viral vector was made into the dorsal cochlear nucleus, followed by an incubation period of two to four weeks. Subjects were then reanesthetized, and blue light (480nm) was delivered via flexible optical fiber to the surface of the CN. Auditory brainstem responses (ABR) were measured and multi-channel recordings of neural activity were made in inferior colliculus (IC) and auditory cortex (Actx) during acoustic, electrical, and light stimulation. The presence of ChR2-expression in CN neurons was confirmed with post-experiment histology.

Results:
Light-mediated responses were recorded from the IC during pulsed blue light exposure, correlating with ChR2 expression within the CN. Responses increased with elevated radiant levels and diminished when the optical fiber was directed away from the CN. Responses evoked with stimulation using a large-diameter (400 um) fiber showed a broad pattern of excitation in IC similar to electric stimulation. The extent of excitation and threshold of responses did not appear to depend on stimulation rate. An increase in neural activation of auditory cortical neurons was also observed during optical stimulation. No significant increase in neural activity was measured in cases where ChR2 expression was outside the auditory brainstem and in control cases. Finally, there was no measurable auditory artifact while using the blue light laser.

Conclusions:
Neurons expressing AAV-ChR2 in the CN can mediate light activation of auditory pathways in the midbrain and cortex in vivo. This approach has the potential to offer more selective stimulation of limited regions of the CN by delivering precise light stimulation via light-emitting diode-based optrodes.

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Infrared Neural Stimulation (INS), an Alternative to Neural Interfaces
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Background: Infrared neural stimulation describes a method, by which an infrared laser is used to stimulate neurons, their axons, or their dendrites. Pulses of mid-infrared laser radiation are delivered to native, non-modified neural tissue to depolarize the neurons. The main advantage of INS is the possibility of stimulating neuron populations with a high spatial resolution when compared with electrical stimulation. Implantation of a miniaturized light source into the cat cochlea demonstrated that stimulation evoked a behavioral response of the animal and showed that stimulation over weeks of time was possible. At the wavelengths used for INS, the energy is primarily absorbed by water in the tissue and is converted into heat. Recently, it has been demonstrated that local heating of the neural tissue by the laser pulse induces a capacitive and depolarizing current at the cell membrane. To translate the technique into a clinical application it is important to know the energy required to stimulate the neural structure. With this study we provide measurements of the radiant energy, radiant power, and radiant exposure at the target structure that is required to stimulate the auditory neurons.

Methods: Flat polished fibers were inserted into scala tympani that the spiral ganglion was in front of the optical fiber. Angle polished fibers were inserted along scala tympani and the beveled surface of the fiber allowed directing the radiation beam perpendicular to the spiral ganglion by rotating the fiber.

Results: The radiant exposure for stimulation at the modiolus for flat and angle polished fibers was on average 6.78±2.15 mJ/cm². With the angle polished fibers, changing the orientation of the optical beam by ~90° from an orientation that resulted in a maximum INS-evoked response, decreased the response amplitude by about 50%. An orientation of the beam that was opposite to the orientation with the maxima, resulted in a minimum.

Conclusions: Optical stimulation is possible with flat polished and side firing optical fibers. When compared with electrical stimulation, an optical implant would require similar amounts of energy to simulate the neurons.

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Optogenetic Control of Mouse Outer Hair Cells

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**Background:** Normal hearing in mammals depends upon both sound detection by inner hair cells (IHCs) and sound amplification by outer hair cells (OHCs). The somatic motility of OHCs driven by membrane potential changes is a major component of the cochlear amplification process. A variety of methods have been utilized to study OHC motility. However, all these techniques have considerable limitations such as being invasive and having a lack of cellular precision and temporal-spatial resolution. To overcome these limitations, we used an optogenetic approach based on channelrhodopsin-2 (ChR2), a direct light-gated non-selective cation channel (NSCC) derived from the blue-green algae *chlamydomonas reinhardtii*.

**Methods:** Patch-clamp and whole cell recordings were performed on the auditory cell line, HEI-OC1, and on isolated mouse OHCs expressing ChR2-tomato. For in vivo analysis, ChR2(H134R)-tdTomato was conditionally expressed in OHCs under control of a prestin-CreER12 transgene. Auditory Brain Stem responses and histological analysis were conducted on these mice.

**Results:** We showed that 100 ms light pulses (λ = 473 nm, 1 mW/mm2) elicited a typical ChR2 current in ChR2(+) HEI-OC1 cells: an inward current rapidly rising to a peak (~15.9±2.1 pA/pF at ~60 mV), and then relaxing to a sustained current with reduced amplitude in solutions blocking all K channels. The I/V plot showed a reverse potential (Vr) near 0 mV. With regular Na+ rich extracellular and K+ rich intracellular solutions, similar ChR2 currents were elicited with the Vr at ~0 mV. I-clamp mode showed that the light pulse transiently depolarized ChR2(+) HEI-OC1 cells by ~11.7 mV. In comparison, in ChR2(-) HEI-OC1, the light elicited no response in membrane current or potential. In isolated mouse ChR2(+) OHCs, similar ChR2 currents were elicited with ~-4.0 pA/pF at ~60 mV in solutions blocking K conductances, with I-V plots showing Vr at ~0 mV. In isolated ChR2(-) OHCs, the light elicited no current or potential change. Analysis of adult (8 weeks) mice with OHC-specific ChR2-tomato expression revealed that nearly 100% of OHCs expressed the transgene and possessed normal ABR and DPOAE levels.

**Conclusions:** This is the first demonstration that ChR2 was successfully expressed in mouse OHCs and HEI-OC1 cells and functionally exhibited a light-activated NSCC and depolarization. The data suggests a novel approach to modulate OHC membrane potential, the driving force of transduction current and the regulator of somatic motility andrepresents a step towards a potential non-invasive clinical intervention of cochlear amplification. Supported by NIDCD DC 00105, 004716 and 00141.

Optogenetic Stimulation of the Auditory Nerve

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When hearing fails, electrical stimulation of spiral ganglion neurons (SGNs) by cochlear implants (CIs) enables speech comprehension despite poor frequency and intensity resolution. Optical SGN stimulation may improve CI coding by enabling more independent information channels to the CNS. We employed the light-gated ion channel, channelrhodopsin-2 (ChR2), to render SGNs sensitive to light. Stimulation of ChR2-expressing SGNs by blue light activated the auditory pathway in hearing mice and in deaf mice. ChR2-mediated auditory brainstem responses (oABR) had a threshold of 2 μJ/mm2 and a dynamic range of more than 20 dB. In summary, optogenetic stimulation of SGNs is feasible.
The Role of Actomyosin Contractility in PCP Regulation in the Organ of Corti
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Background
Planar cell polarity (PCP) in the organ of Corti (OC) is manifested by the uniform orientation of stereociliary bundles towards the lateral edge of the cochlear duct. Proper orientation of stereociliary bundles is essential for normal hearing function as they are directionally sensitive to vibrations along the basilar membrane. In addition to the core cassette of non-canonical Wnt/PCP pathway genes, Protein tyrosine kinase 7 (Ptk7) was identified as a novel regulator of PCP in vertebrates. In Ptk7 mutant mice, outer hair cells (OHCs) display stereociliary bundle misorientation along with extra rows of OHCs in the apex, similar to core PCP mutants.

Methods
We hypothesize that Ptk7 and the core Wnt/PCP pathway converge on myosin II to mediate PCP in the organ of Corti. Myosin II-based contractile tension between supporting cells and hair cells orient hair cell PCP. To test this hypothesis, we are taking a multipronged approach combining mouse genetics, organotypic cochlear explant culture, small molecule inhibitors and live cell imaging to elucidate the role of the actomyosin cytoskeleton in hair cell PCP.

Results
We show that Ptk7 and the core Wnt/PCP pathway differentially regulate a contractile myosin II network in supporting cells. Moreover, Ptk7 and the Wnt/PCP pathway act in concert to promote junctional planar asymmetry of vinculin, a tension-sensitive actin-binding molecule. To gain mechanistic insight into this process, we are currently analyzing the dynamics of myosin II during PCP signaling in the organ of Corti by live imaging of cochlear explants expressing a GFP-Myosin IIB fusion, and assessing the effect of perturbing the actomyosin cytoskeleton on contractile behaviors of myosin IIB. We also generated conditional vinculin knockout mutants and observed PCP defects in the vinculin mutant inner ear. These results suggest a model whereby PCP signaling mediates polarized contractile forces along the medial borders of hair cells to orient their stereociliary bundles. Details of our analysis will be presented at the meeting.

Conclusion
Our results support the hypothesis that Ptk7 and the non-canonical Wnt/PCP pathway act in concert to mediate myosin II-based anisotropic tension at contacts between hair cells and supporting cells. We showed previously that hair cell PCP is controlled by Rac-PAK signaling. We propose that polarized contractile tension between OC epithelial cells regulates the spatial pattern of Rac-PAK activity to orient hair cell PCP.
Patterning of the organ of Corti is a multi-step process that involves dual modes of Notch signaling

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Background
Notch-mediated lateral inhibition is a well-characterized patterning mechanism in the organ of Corti; however, it is insufficient to explain the intricate array of hair cells and supporting cells in the mammalian cochlea. Here we propose a dual mode of Notch action that is dose-dependent and acts differentially on progenitors that generate inner and outer hair cells.

Methods
We compared several mutants of the Notch pathway in the inner ear, analyzing the appearance of supernumerary hair cells and whether or not they were accompanied by transdifferentiation of associated supporting cells. We are also pharmacologically manipulating levels of Notch signaling in organotypic cochlear cultures to evaluate a dose dependency effect.

Results
In inner ear conditional mutants where some components of the Notch signaling pathway have been deleted, we do not observe supernumerary outer hair cells nor transdifferentiation of supporting cells to a hair cell fate. By contrast, we do observe supernumerary inner hair cells and their associated supporting cells.

Conclusion
Our results so far support a model in which low levels of Notch signaling might be sufficient to pattern inner hair cells, border cells and inner phalangeal cells whereas higher levels of Notch may be required to correctly regulate the proportions of outer hair cells and pillar and Deiters’ cells.

Vangl2 Directs Supporting Cell Morphogenesis and is not Essential for the Postnatal Refinement of Hair Cell Planar Polarity

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Background
Auditory hair cells have a distinctive planar polarity evident in the polarization and orientation of the stereocilia bundle. Mutations in the core planar cell polarity (PCP) gene vangl2 result in hair cells that form polarized bundles but are misoriented and have bundle polarities that are not coordinated with neighboring cells. In addition there is a gradient of phenotypic severity ranging from the least affected basal to the most affected apical turn of the cochlea. This gradient is similar to the progression of hair cell maturation suggesting that active refinement could correct planar polarity phenotypes in the basal cochlea of vangl2 knockout (KO) mice.

Methods
Since the vangl2 deletion results in perinatal lethality, vangl2 conditional knockouts (CKO) were generated to test this hypothesis. When crossed with pax2-Cre, vangl2 is deleted from the inner ear yielding planar polarity phenotypes similar to vangl2 KOs at late embryonic stages. Early postnatal development was assayed using anatomical techniques and auditory function in adults was assayed by ABR and DPOAE.

Results
Pax2-Cre; vangl2 CKO mice are viable and do not have the lethal neural tube defect that occurs in the vangl2 KO. Quantification of planar polarity through postnatal development demonstrates a continuation of the maturational refinement processes initiated during late embryogenesis, and auditory hair cell orientation is largely corrected within 10 days of birth although some misoriented cells persist in the extreme apical turn of the cochlea. In contrast a similar refinement of vestibular hair cell planar polarity in the striola region of the utricular maculae does not occur during this period. In addition the pax2-Cre; vangl2 CKO has profound changes in the shape and distribution of the apical surfaces of Outer Pillar Cells and Deiters’ Cells. These changes in supporting cell morphology are persistent, and are not corrected during the period of planar polarity refinement. ABR analyses of adults show a 10-20 decibel shift in auditory
threshold across all tested frequencies and DPOAE measurements indicate that this mild hearing deficit is of cochlear origin.

Conclusion
Together these data support the hypothesis that a vangl2-independent refinement mechanism actively reorients auditory hair cells during the first 10 days of postnatal development. Vangl2 is required during supporting cell development however it is unclear if this reflects a ‘conventional’ planar polarity activity. Nonetheless since the altered supporting cell morphology is not refined these changes in supporting cell shape likely underlie the hearing deficits measured in vangl2 CKOs.

GSK3 is an essential regulator of cell fate decisions in the mouse cochlea
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Background
The development of the cochlear sensory epithelium requires the orchestration of multiple developmental events including cell cycle-exit, precise cell fate specification, and organized alignment of hair cells and supporting cells. Many of the signaling pathways required to coordinate these processes remain unknown. In the central nervous system the cytoplasmic molecule Glycogen Synthase Kinase 3 (GSK3) acts as an integrating molecule for multiple proliferation and differentiation signals. In particular, GSK3 is regulated by receptor tyrosine kinase, canonical Wnt, and Sonic hedgehog signaling pathways. While canonical Wnt signaling has been implicated in auditory versus vestibular fate specification, prosensory formation and hair cell differentiation, functional role(s) for GSK3 in the developing cochlea have not been examined.

Methods
To begin to identify the functional role(s) for GSK3 signaling in cochlear development, we established explant cultures from E13.5 mouse embryos. GSK3 was inhibited using one of two different pharmacological inhibitors (BIO-acetoxime and CHIR99021).

Results
Inhibition of GSK3 leads to several unique phenotypes including a disruption in hair cell arrangement, an increase in the number of inner hair cells (confirmed based on expression of Fgf8), and a widened gap between the row of inner hair cells and the first row of outer hair cells. This phenotype was markedly different from the phenotype observed in explants treated with the canonical Wnt agonist, suggesting that the effects of GSK antagonism are not a result of activation of canonical Wnt signaling. Inhibition of GSK3 also leads to a decrease in expression of Bmp4 in cochlear explants, suggesting that BMP may be a target of GSK3.

Conclusion
Our data suggest that non-Wnt-related GSK3 signaling plays an important role in cochlear development that may include regulation of inner and outer hair cell fates.

The Role of Atoh1 in the Cochlear Development
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Background
Atoh1, the mouse homolog of Drosophila proneural gene atonal, is a basic helix-loop-helix (bHLH) transcription factor that is the earliest known gene expressed in differentiating hair cells. There is a severe phenotype in the cochlea of Atoh1-null mice: massive cell death in the presumptive sensory epithelia begins at embryonic day 15.5 (E15.5) and results in the complete absence of hair cells and supporting cells, suggesting the necessity of Atoh1 in the development of the organ of Corti.

Methods
To dissect Atoh1’s function during hair cell development, we established a conditional knockout (CKO) system that allows us to delete Atoh1 from hair cells at different stages. By exposing pregnant or neonatal mice to tamoxifen to activate Cre-
mediated recombination in hair cells, Atoh1 was removed from hair cells at stages between E15.5 and postnatal day 0 (P0).

Results
We analyzed Atoh1 protein expression in mice in which both copies of the Atoh1 gene have been replaced by an Atoh1-GFP fusion sequence. Based on our results of GFP immunohistochemistry and quantitative PCR in these mice, Atoh1 expression initiates in the prosensory domain of the cochlea as early as E13.5. As the cochlea develops, Atoh1 expression is up-regulated and maintained in hair cells until postnatal stages, when it starts to be down-regulated and disappears from hair cells before the onset of hearing.
We analyzed the number of hair cells and supporting cells in the cochleas of the Atoh1-CKO animals and found there is a critical time window (about two days after Atoh1 expression) in which Atoh1 is required to maintain the survival of the cells in the organ of Corti. We also examined the hair bundles in the CKO hair cells and found the hair bundle formation also requires Atoh1 expression in a time-dependent manner. In addition, auditory brainstem response (ABR) was measured in the CKO animals in which Atoh1 was removed at E17.5. Although we observed comparably normal hair cell formation in the basal region of the cochlea in these mice at neonatal stages, we could not detect ABR responses at any frequency in these mice at the age of 6 weeks, indicating an essential role of Atoh1 in maintaining the proper function of hair cells after their differentiation.

Conclusion
Our study suggests Atoh1 might have multiple functions in the survival, differentiation and maturation of hair cells during cochlear development.
Using Novel Mouse Mutants to Elucidate bHLH Transcription Factor Regulatory Interactions in Ear Neurosensory Development
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Background
Cross-regulation between transcription factors plays crucial roles in cell differentiation and development. In developing ears, studies have shown extensive intra- and intercellular interactions among the three proneural basic helix-loop-helix (bHLH) transcription factors, Neurog1, Neurod1 and Atoh1. Together with interactions with a group of diffusible factors and other transcription factors, they form a regulatory network that determines neurosensory cell identity.

Methods
We generated several novel mouse mutants and characterized their inner ear neurosensory development and gene expression using immunohistochemistry and in situ hybridization.

Results
Using Neurod1 conditional knockout mutant mice, we show that the absence of Neurod1 results in upregulation of Atoh1 expression and hair cell formation within the sensory ganglion, suggesting that Neurod1 negatively regulates Atoh1 to suppress hair cell fate in neurosensory precursor cells to develop as neurons. Moreover, the deletion of Neurod1 also results in premature and increased expression of Atoh1 and formation of ectopic inner hair cells in the outer hair cell region in the apex of the organ of Corti. Interestingly, the phenotype of ectopic inner hair cell formation can be partially suppressed by deleting one copy of Atoh1 gene, suggesting that the expression level of Atoh1 may control type-specific hair cell differentiation. To further investigate the cross-interactions between Atoh1 and Neurod1, we combined the deletion of Neurod1 with the Atoh1 ‘self-terminating’ conditional knockout, in which the Cre expression is driven by an Atoh1 enhancer element and activated when Atoh1 protein binds. In addition to the Atoh1 self-regulatory binding site, the enhancer element also contains a putative Neurod1 binding site. The resultant mutant mice showed increased hair cell loss throughout the cochlea compared with the Atoh1 ‘self-terminating’ mutant. This suggests that Atoh1 expression may be more rapidly reduced due to increased Cre expression, which is possibly the combined effect of both the direct elimination of Neurod1 inhibition on Cre expression and the indirect increased Atoh1 self-promoting activity.

Conclusion
Our results suggest that the cross-interactions between Neurod1 and Atoh1 may provide molecular basis of neurosensory cell fate determination and type-specific hair cell differentiation. We will quantitatively measure and compare the expression level of relevant genes in these different mutant ears. Our data will help to further define the bHLH transcription factor regulatory network and the role of Atoh1 expression level and duration in hair cell differentiation and maintenance, which will provide crucial information for reconstituting the organ of Corti.
Function and regulation of Hey1 and Hey2 in cochlea development
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Background
The mouse auditory sensory epithelium consists of mechano-sensory hair cells (HCs), critical for our ability to detect sound, and supporting cells (SCs). Both cell types derive from a common pro-sensory progenitor. At E14.0, shortly after terminal mitosis HC and SC differentiation initiates in a basal to apical gradient. Very little is known about the molecular mechanisms controlling the timing of HC and SC differentiation. Hey1 and Hey2, two highly redundant bHLH transcriptional repressors are highly expressed in cochlear pro-sensory progenitors and are rapidly down-regulated as differentiation occurs.

1. Based on their known function as Atoh1 antagonists and their high expression in pro-sensory progenitors, we hypothesize that Hey1 and Hey2 function in maintaining an undifferentiated progenitor state.
2. Hey1 and Hey2 have been commonly described as Notch effector genes. However, our recent findings suggest a more complex regulation mechanisms for Hey2 during cochlear development. A good candidate pathway to regulate Hey2 in pro-sensory progenitors is the Sonic hedgehog (Shh) pathway, which has been shown to play an inhibitory role in HC differentiation.

Methods
1. Pax2-Cre Hey1 fx/fx Hey2 null mice were used to genetically delete both Hey1 and Hey2 in the developing cochlea.
2. We cultured E13.5 Atoh1/GFP transgenic cochlea explants in the presence of Shh and cyclopamine and vehicle control and used qpcr to analyze changes in gene expression. In addition effects on HC differentiation were analyzed over a three day period monitoring HC specific Atoh1/GFP expression.

Results
1. Consistent with abnormal timing of differentiation, Hey1 Hey2 double knockout mice have complex hair cell patterning defects. We observe in the Hey1 Hey2 double mutant sensory epithelium ectopic IHCs from base to apex and ectopic OHCs in the base and mid cochlear regions. However, we observe in the apex of the Hey1 Hey2 mutant cochlear only two rows of OHCs.
2. Our results indicate that Shh positively regulates Hey2 expression at the pro-sensory stage. Moreover, cochlea explants treated with Shh show a delay in HC differentiation, while in those treated with the Shh antagonist, Cyclopamine, HC differentiation occurs earlier.

Conclusion
1. Our results show a patterning defect that suggests an early onset of HC in absence of Hey1 and Hey2 in the double knockout mutant mice possibly leading the altered phenotype.
2. Based on our findings we hypothesize that Shh pathway negatively controls HC differentiation through a Hey2 dependent mechanism.

Maintenance of cell fate in the organ of Corti: molecular biology of Atoh1 regulation by Hes/Hey genes
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Background
A regular mosaic of hair cells and supporting cells characterizes the sensory regions of the inner ear. Genetic experiments indicate that this mosaic arises partly through Notch-mediated lateral inhibition leading to regulation of Atoh1 gene expression. During embryonic development, Atoh1 up-regulation is required for hair cell differentiation of selected sensory progenitors, and coincident inhibition in the surrounding progenitors, which subsequently differentiate as supporting cells. A requirement for Notch signaling continues to be required during perinatal maturation of the organ of Corti to maintain the fate of supporting cells. The molecular basis of Atoh1 gene regulation during these processes remains to be studied.

We have investigated the role of the Hes/Hey family of transcriptional repressors in the suppression of Atoh1 expression, and thus the maintenance of supporting cell fate. Hes/Hey expression in most perinatal supporting cells depends on the continued activation of the Notch signaling pathway. Blocking Notch signaling leads to Atoh1 activation and supporting cell transdifferentiation. Our working hypothesis is that Atoh1 is actively suppressed in supporting cells by
Hes/Hey factors. In response to the loss of Notch activity in supporting cells, Atoh1 up-regulation is mediated by a process of disinhibition caused by the loss of Hes/Hey activity. Experiments supporting this hypothesis will be described.

Methods
Chromatin immunoprecipitation (ChIP), organ culture, FACS purification of hair cells and supporting cells, in vitro transfection, site-directed mutagenesis, and molecular reporter assays were used to investigate the role of Hes/Hey genes in regulation of Atoh1 expression.

Results
Hes/Hey gene expression is rapidly downregulated when Notch signaling is blocked, and Atoh1 is rapidly up-regulated. ChIP assays provide evidence of direct binding of Hes/Hey proteins to the Atoh1 gene in a model cell line. Transfection assays indicate that Hes/Hey are sufficient to bring about Atoh1 transcriptional down-regulation, and that this likely occurs through direct binding to Atoh1 gene regulatory regions, and the changes in epigenetic regulation that accompany this binding.

Conclusion
Hes/Hey genes contribute to the regulation of Atoh1 through a mechanism that involves direct binding of these transcription factors to Atoh1 gene regulatory elements. Regulation of Atoh1, Hes/Hey genes and the importance of these observations to both the maintenance of the differentiated state, and regeneration, will be discussed.

Tinnitus specific trait in animals
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1
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Tinnitus is a non-curable stress-related brain disorder, that is mostly noise-induced and whose origin is unknown. We have addressed the molecular and physiological basis of this disease using a combined approach that included behaviorally tested tinnitus (Rüttiger et al., Knipper, Hear Res 2003), hearing measurements (including DPOAEs, ABRs and ABR wave analysis) and markers that trace network activity (Arc/Arg3.1). Data analysed the first time equally hearing impaired animals that were behaviorally distinguished in hearing impaired animals with and without tinnitus. We compared animals between the periphery of the cochlea and the auditory cortex, including the hippocampus and amygdala. We also included an analysis of altered responsiveness after stress priming. We unraveled a tinnitus specific trait that may explain some of the existing controversies about the molecular basis of tinnitus.
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Short-term traumatic and long-term non-traumatic noise exposure both increase spontaneous firing rate and synchrony in cat auditory cortex
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1
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Following traumatic noise exposure, resulting in a permanent hearing loss, the primary auditory cortex characteristic frequency (FF)-place map becomes reorganized. Instead of representing the frequency region of the hearing loss, now those neurons are tuned to the edge frequency of the audiogram, resulting in an over representation thereof. For this reorganized cortical region, spontaneous firing rates (SFR) are approximately doubled and spontaneous neural firings are occurring in a more synchronized way. Following frequency-restricted long-term moderate level (<70 dB SPL) stimulation with noise or other multi-frequency sounds, no changes in ABR thresholds are found and DPOAEs remain normal. Yet, in auditory cortex the region with CFs in the frequency range of the sound exposure becomes much less responsive, it's SFR decrease as well as the slope of the driven-rate-level functions. In contrast, on both the low- and high-frequency side, the cortical gain is increased leading to increased SFR, increased neural synchrony and increased slope of the driven-rate-level functions. These changes require up to two weeks to establish and more than 3 months for recovery in quiet. Important differences are that the increased SFR following noise trauma is not accompanied by increased cortical gain, whereas this is the case following the long-term moderate level exposure. Following traumatic noise exposure, tinnitus is a frequent finding, and from the results induced by long-term moderate level noise one would expect the same, i.e., tinnitus but in the absence of hearing loss or peripheral auditory damage, but likely accompanied by hyperacusis.
Psychoacoustic and Electrophysiological Imaging of Tinnitus
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BACKGROUND: Neural mechanisms that detect changes in the auditory environment rely on processes that predict sensory state. This principle suggests that in tinnitus there is a disparity between what the brain predicts it is hearing (this prediction coded by aberrant neural activity occurring in frequency regions affected by hearing loss) and the acoustic information delivered to the brain by the damaged cochlea. This disparity may activate auditory attention, facilitating through subcortical neuromodulatory systems forms of neural plasticity that entrench aberrant neural changes underlying the tinnitus sound. This talk will summarize behavioral and functional brain imaging evidence for persisting auditory attention in tinnitus and present a qualitative model describing how attention may operate in normal hearing and in tinnitus.

METHODS: We measured hearing function, tinnitus spectra, and residual inhibition (RI) functions in 28 individuals with tinnitus, and hearing function in 31 age-matched controls without tinnitus. Audiometric thresholds to 16 kHz were comparable in the two groups. We then probed neural activity with 40-Hz amplitude modulated tones using either a 500 Hz carrier frequency in the range of normal hearing or a 5000 Hz carrier in the tinnitus spectrum and the region of audiometric threshold shift. Each subject adjusted the loudness of the probe stimulus to match a 1000 Hz sound (in the range of normal hearing) presented at 65 dB SL. This procedure may control for changes in central gain related to hearing loss and reveal consistent electrophysiological findings presently lacking in the tinnitus literature. Two responses were extracted from 128 channel EEG, the 40-Hz steady state response (ASSR) localizing to A1 and the N1 transient response localizing to A2, both responses known to be attention sensitive. The responses were also measured after a masking sound (band limited noise, center frequency 5000 Hz) presented for 30 seconds that yielded RI in tinnitus.

RESULTS: N1 amplitude was larger at both probe frequencies and ASSR amplitude at 500 Hz in the tinnitus group compared to controls. At 5000 Hz ASSR amplitude was smaller in tinnitus than controls. The group difference in N1 was not affected by masking, but the group difference in ASSR amplitude was abolished by masking.

CONCLUSIONS: These results can be interpreted assuming that (1) frequency-nonspecific auditory attention is activated by a failure of auditory prediction in tinnitus, and (2) neuroplastic changes occurring in the tinnitus frequency region bind neurons into synchronous activity underlying the tinnitus percept.

Neuroimaging of tinnitus-related changes in the human central auditory system
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Tinnitus is associated with peripheral hearing loss in the majority of cases. However, only a minority of people with a peripheral hearing loss experiences tinnitus. Evidently, peripheral hearing loss is not a sufficient condition to develop tinnitus, and brain-related factors have been hypothesized to play a role in the development of tinnitus. We applied functional MRI to investigate brain mechanisms involved in tinnitus.

One specific hypothesis that we tested is that tinnitus is related or even caused by reorganization of the tonotopy in the auditory cortex. However, an fMRI study in subjects with tinnitus and (near) normal hearing did not show reorganization. A second hypothesis, that assumes that tinnitus is related to reduced inhibition, predicts that sound-evoked responses may be enhanced in tinnitus. This hypothesis was tested by a comparison of subjects with and without tinnitus, respectively, and matched moderate hearing loss. There was no clear evidence of enhanced responses in the tinnitus subject group, which did not support the hypothesis. An additional analysis investigated the correlation between activity in auditory structures in the brainstem, thalamus and cortex. As was to be expected, the brainstem and cortex each have a strong functional connection with the thalamus. However, the functional connectivity between the brainstem and cortex was reduced in tinnitus subjects. This reduced connectivity may be an indicator of abnormal gating of auditory information in the thalamus. A third experiment was performed in subjects with gaze-modulated tinnitus. These subjects all underwent acoustic schwannoma surgery. They were selected for the ability to modulate the loudness of their tinnitus by peripheral gaze. An increase of the tinnitus loudness was associated with an increase of activity in the brainstem and a reduction of inhibition in the cortex. Furthermore, lateral gaze was associated with inhibition of activity in the thalamus. This unexpected inhibition again suggests abnormal thalamus function in tinnitus.

Together, these data show that tinnitus does not require reorganization of the cortical tonotopic map. Rather, functional correlates of tinnitus suggest a pivotal role of the thalamus in tinnitus. These correlates could either be a consequence of
tinnitus, or may reflect a pre-existing vulnerability to develop tinnitus. Also, they could arise as a consequence of inherent mechanisms in the central auditory system or could be induced by interactions with non-auditory structures. Future research must help to understand these causal relations, which may help to develop new therapies for tinnitus.

**Stress and Tinnitus: Population and experimental data**

_Sylvie Hebert_¹,², Dan Hasson³

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**Title**: Stress and tinnitus: Population and experimental data

Sylvie Hébert
Dan Hasson

Hearing problems are a public health issue with prevalence figures far more common than previously estimated. There are well-established risk factors of hearing problems such as age, sex and noise exposure history. Our population studies systematically demonstrate an association between symptoms of long-term stress and increased prevalence of hearing loss and tinnitus. In particular, emotional exhaustion has been found a stronger predictor of both tinnitus presence and severity than traditional risk factors, e.g. noise at work, sex and hearing loss. Our experimental studies involving cortisol analyses, a stress-related hormone, also support an association between tinnitus and long-term stress. In particular, in tinnitus there is an increased sensitivity of the hypothalamus-pituitary-adrenal (HPA) axis to negative feedback. Such HPA axis disturbance has been described in other clinical populations such as patients with chronic fatigue syndrome and burnout. Altogether these findings have implications on treatment options for individuals suffering from severe tinnitus and suggest that measures of stress need to be taken into account in order to proactively intervene and prevent hearing problems. In this session, we will summarize our recent population and experimental findings about the associations between stress and hearing problems, and more specifically tinnitus.

**Light on network dynamics in primary auditory cortex**

_Israel Nelken_¹

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The last few years have seen the introduction of a number of exciting new techniques to auditory neuroscience, including calcium imaging of large populations of neurons and optogenetic techniques for the identification of neurons and manipulation of their responses. In this talk, I will highlight one application of these techniques - the study of the correlation structure of large neuronal ensembles, and its modification by experience. Auditory cortex is malleable by experience, but previous studies of experience-dependent changes in auditory cortex emphasized modifications at the single-neuron level or modifications of the tonotopic map. Here, I will describe a study of the influence of a dramatic yet natural experience in the life of female mice - giving birth and becoming a mother - on neuronal ensembles in primary auditory cortex. Groups of neurons in layers 2/3 of mothers and age-matched controls were monitored using in-vivo two photon calcium imaging, verified and complemented by electrophysiological recordings from single neurons. Stimuli consisted of a set of artificial sounds and natural pup vocalizations. While response profiles of single neurons showed subtle changes in mothers relative to non-mother controls, population dynamics underwent large changes as measured by a large increase in pairwise and higher-order noise correlations: pairwise noise correlations essentially double in size in mothers relative to non-mother controls, independently of the tested stimulus. We used decoding techniques to estimate the influence of the increased noise correlations on stimulus coding. Surprisingly, decoding performance remained essentially the same in the two groups and was not affected by removing noise correlations from the responses. Thus, the correlated activity of the cortical network appears to be essentially orthogonal to its information-carrying capabilities.

**Two-photon imaging of neuronal activity in the mouse auditory cortex reveals spontaneous categorization of sounds**

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In categorical perception a set of smoothly changing stimuli is perceived in a discontinuous manner. The neuronal correlates of this phenomenon are poorly understood. Here, we used in vivo two-photon calcium imaging to analyze sound-evoked activity patterns in local neuronal ensembles of the mouse auditory cortex. For that we used a large library
of ~70 different sound stimuli containing pure-tone pips and various complex sounds. We found that activity patterns are highly constrained into few discrete response modes, which is surprising given how many patterns could theoretically be generated by the combination of already a few neurons. Such a low number of observed response modes suggest that local activity patterns form a discrete representation of sounds. Using on-line synthesis of sounds to tailor stimuli that allow precise probing of a given neuronal population, we observe highly non-linear dynamics indicating an antagonistic ‘winner-takes-all’-like competition between response patterns. The high constrain on patterns that can be evoked implies that the information encoded locally is very low. However, we observed that different combinations of sounds were associated in a response mode in different local populations so that combining the information of multiple local populations forms a representation that allows discrimination of more than 80% of sound pairs on a single trial basis. To test if discrete representations of sounds in the auditory cortex reflect auditory percepts in the mouse we trained mice in a go/no-go task to discriminate a positively reinforced sound and a negatively reinforced sound. We here used a novel assay where the spontaneous responses of mice to non-reinforced off-target sounds reported the perceived similarity of many off-target sounds to the reinforced target sounds. We observed non-linear categorization behavior in response to linear morphings of two target sounds. Interestingly, we observed a good match of the prediction based on global activity patterns recorded in the imaging experiments and behaviorally measured perception, including its non-linearities. In summary, our results suggest that local non-linear dynamics shape the cortical representation of sounds into a basis set of spontaneous categories that are available for behavioral decisions.

Micro-organization and plasticity of the primary auditory cortex
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The auditory cortex is a laminated structure organized into radial columns that adaptively processes sensory information from the external environment. The precise nature of the transformation of sensory information at the level of cortical networks is unknown. We use in vivo two-photon calcium imaging techniques to measure response properties and functional organization of primary auditory cortex neurons in mouse. We find that frequency selectivity in supragranular layers is heterogeneous on small spatial scales. In contrast to the supragranular layers, heterogeneity of frequency selectivity is lower in thalamorecipient layers indicating a rapid laminar transformation of the spatial representation of frequency selectivity. These findings reveal a transformation of sensory representations that occurs across a single cortical layer within the auditory cortex, which could generate sequentially more complex analysis of the acoustic scene incorporating a broad range of spectro-temporal sound features. The local spatial heterogeneity of frequency selectivity in supragranular layers is likely created by sampling of diverse inputs to supragranular neurons by intra- and inter-laminar connections. The large frequency range of inputs available to each neuron might provide a substrate for a large degree of plasticity in individual neurons. We tested the capacity of auditory cortex neurons to rapidly change tuning properties by using micro-stimulation of top-down projections to auditory cortex and pairing such stimulation with a particular sound. We find that the frequency tuning of individual neurons can rapidly be changed leading to an increase in the representation of the paired sound. Collectively, these results provide insight into how sensory information is represented and adaptively transformed in auditory cortex.

In vivo two-photon calcium imaging of dendritic spines in auditory cortex
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The dendritic spines are key structures for receiving excitatory inputs and for synaptic plasticity. Understanding the individual functional properties and spatial arrangement of afferent synaptic inputs on dendrites is essential to dissect how the information is processed by neurons in the mammalian brain. However, the small size and high density have precluded the functional study of dendritic spines in vivo with standard methods. We have recently established a new variant of high-resolution two-photon imaging (LOw-power Temporal OverSampling, LOTOS) to detect calcium transients in single spines of mouse cortical neurons in vivo. Using this technique together with whole-cell patch-clamp recordings, we have studied two questions: 1) how is the spatial arrangement of sound-evoked inputs on dendrites in auditory cortex? 2) how individual synapses support cortical up-states in vivo. For the first question, we find that calcium signals require the activation of NMDA (N-methyl-D-aspartate) receptors. Active spines are widely distributed on basal and apical dendrites and pure-tone stimulation at different frequencies reveals both narrowly and widely tuned spines. Interestingly, spines tuned for different frequencies were highly interspersed on the same dendrites: even neighboring spines were mostly tuned to different frequencies. These results demonstrate that NMDA-receptor-dependent single-spine synaptic inputs to the same dendrite are highly heterogeneous (Chen et al., Nature, 2011). For the second question, we demonstrate that NMDA receptor-dependent subthreshold synaptic spine calcium signals are present during up-states, but nearly absent
during down-states. We estimate that more than 500 excitatory synapses are active during an up-state. Importantly, similar patterns of spine calcium signaling are observed during spontaneous and sensory-evoked activity, indicating that cortical up-states and sensory stimulation-evoked cortical responses may be driven by highly overlapping sets of upstream cortical neurons. These results identify the dendritic pattern of glutamatergic synaptic inputs on the level of single spines during cortical up-states. Altogether, with the high-resolution two-photon imaging technique, we open a way for in vivo functional study of the sensory information processing and the cortical network dynamics at single synapse level.

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Single neuron and local network plasticity – insights from motherhood
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Understanding sound processing in the auditory cortex can benefit from recent imaging methods like two-photon laser scanning microscopy. One advantage of imaging is that it allows measurements at high spatial resolution and from multiple cells simultaneously. I will describe our recent experiments using in vivo two photon imaging to study how pure tones and natural sounds are represented in layer 2/3 neurons in the primary auditory cortex (A1) of virgin vs. primiparous mother mice. To obtain a mechanistic understanding of network plasticity we targeted excitatory and inhibitory neurons selectively. We used either wild type NMRI mice or a cross between a knock-in parvalbumin positive driver mouse with a tdTomato reporter mouse. For calcium imaging we injected Fluo-4AM into L2/3 and imaged the cortex with a Prairie two-photon microscope. Electrophysiology was limited to juxtasomal recording either using blind patch or using visual guidance. Using calcium imaging we found that local populations display low signal correlations but high and variable noise correlations. Using electrophysiology we show that PV neurons provide a unique form of inhibition onto excitatory neurons. Moreover, PV neurons change their inhibitory pattern onto excitatory cells following the transition to motherhood and when multisensory inputs are processed. Our data show that plasticity in mothers is tunneled via modulation of feedforward inhibition.

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Corticostriatal Projections Mediate Auditory Decisions
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How does the brain use auditory signals to guide behavior? Representations of the acoustic world are constructed in the auditory cortex (ACx), but how these representations are used to effect behavioral choices is largely unknown. Auditory information is relayed from the ACx to a number of cortical and subcortical targets by distinct, largely non-overlapping populations of pyramidal neurons. We hypothesized that the ACx drives subjects’ choices through its projection to the striatum. To test this hypothesis, we measured the contribution of striatal projection neurons to behavioral decisions by specifically manipulating their activity in rats performing auditory discrimination. We find that the projection of the ACx to the striatum drives auditory decisions in rats.

We first developed a novel auditory task—inspired by classic experiments by Newsome and colleagues in which a random dot stimulus was used to study the representation of visual motion in cortical area MT of macaque monkeys—designed to exploit the tonotopic organization of the ACx. We trained rats to discriminate low- and high-frequency “cloud-of-tones” stimuli (overlapping 30msec pure tones distributed over three octaves) in a two-alternative choice task. Subjects were required to report whether low or high tones were overrepresented; performance varied smoothly with stimulus frequency.

To test the ability of corticostriatal neurons to influence auditory discrimination in this task, we specifically targeted the expression of Channelrhodopsin-2 (ChR2) or Archaerhodopsin-3 (Arch). We then implanted optical fibers into the ACx and limited light intensity to limit the perturbation to a restricted frequency band.

Activation of corticostriatal neurons by ChR2 biased the subjects to select the choice port associated with the neurons’ frequency tuning. Furthermore, inactivation of corticostriatal neurons by Arch made subjects less likely to select the choice port associated with the preferred frequency of the inactivated neurons.

These results show not only that the ACx is causally involved in this task, but also that information propagates beyond the ACx via the corticostriatal projection. As striatal projections are widespread in cortex, our results may point to a general mechanism through which sensory cortex influences decisions.
THE CHROMATIN REMODELING PROTEIN CHD7, MUTATED IN CHARGE SYNDROME, IS REQUIRED FOR PROPER HAIR CELL DEVELOPMENT AND COCHLEAR INNERVATION

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Background

CHD7 is a chromatin remodeling protein that is necessary for proper formation of the mammalian inner ear. Humans with CHARGE Syndrome and CHD7 heterozygous mutations exhibit mixed sensorineural/conductive hearing loss and inner ear dysplasia, including abnormalities of the semicircular canals and Mondini malformations. Prior studies in the mouse have demonstrated that reduced Chd7 dosage in the ear disrupts expression of genes which are critical for morphogenesis and neurogenesis; however, the effects of Chd7 deficiency on cochlear development have not been explored. In mice, complete loss of Chd7 function results in intrauterine death by E11, several days prior to development of the sensory epithelium and its innervation. To overcome this early lethality and to uncover molecular mechanisms of CHD7 function, we have studied Chd7 null phenotypes using inner ear specific Cre and Chd7-flox alleles.

Methods

We performed a detailed analysis of inner ears from mice with FoxG1-Cre-mediated Chd7 conditional deletion, using whole-mount immunofluorescence, paint-filling, and scanning electron microscopy (SEM).

Results

FoxG1-Cre:Chd7 conditional knock-out (CKO) ears have underdeveloped, hypoplastic cochleae, with abnormal rotation of the mid-cochlear region and twisting of the apex that is detectable by E14.5. Myosin-VIIa staining of early postnatal CKO ears shows abnormal rows of outer hair cells along the length of the organ of Corti; at the tip of the apex, hair cells are completely disorganized and do not form typical orderly rows. SEM shows normal stereociliary bundle orientation and supernumerary hair cells throughout the length of the cochlea in CKO mice. Supernumerary hair cell stereocilia appear either cochlear-like or vestibular-like. Innervation of cochlear hair cells in the Chd7 CKO mice is also disrupted, with Neurofilament-positive neurites that extend to the lateral edge of the sensory epithelium. These neurites form loops that project beyond the hair cells, turn and extend backwards toward their site of origin in the cochlear ganglion.

Conclusion

Chd7 is necessary for elongation of the cochlear epithelium, hair cell differentiation and organization, and extension of neurites in the organ of Corti. Phenotypes in Chd7-CKO mice are similar to those observed with disrupted retinoic acid and Notch pathways. We conclude that CHD7-mediated chromatin remodeling regulates early patterning and regulation of hair cell number and differentiation, perhaps via altered retinoic acid and Notch signaling. Further experiments treating Chd7 mutant ears with agents that alter retinoic acid or Notch signaling will explore this possibility.
An analysis of a Sprouty gene dosage series reveals compensatory regulation of the size of the otic placode

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Background

In the mouse, at 8 – 9 somite stages (s), the otic placode becomes morphologically distinct as a pseudostratified region of ectoderm on either side of the hindbrain. These otic placodes are a committed embryonic progenitor pool that contributes to the mature population of mechanosensory hair cells, supporting cells and innervating neurons of the inner ear. We are studying the role of the Sprouty (Spry) gene family (of which there are four murine members, Spry1 – 4) in otic placode induction. Spry genes encode antagonists of signaling downstream of receptor tyrosine kinases, including FGF receptors. In Spry1⁻/⁻; Spry2⁻/⁻ double mutants, the otic placode is enlarged due to an aberrant, increased response of tissue to FGF induction.

Methods

Mutant combinations of Spry1 and Spry2 are generated by crossing β-actin cre/β-actin cre; Spry1⁻/⁻; Spry2⁻/⁻ males to Spry1floxfloxfloxfloxfloxfloxfloxfloxfloxfloxfloxfloxflox females. Embryos are analyzed by in situ hybridization, immunofluorescence, and histology. The size of the otic placode is measured by ImageJ on images of serial sagittal sections through the otic placode. Cell proliferation is quantified as the total number of cells stained for phospho-Histone-H3 per volume of otic placode.

Results

By analysis of a Sprouty gene dosage series, we find that similar to Spry1⁻/⁻; Spry2⁻/⁻ double mutants, the otic placode is also enlarged in Spry1⁻/⁻; Spry2⁻/⁻ and Spry1⁻/⁻; Spry2⁻/⁻ mutant embryos at 9 – 11 s, a stage at which cells have committed to an otic fate. Interestingly, unlike Spry1⁻/⁻; Spry2⁻/⁻ double mutants, enlargement of the otic placode is not maintained in Spry1⁻/⁻; Spry2⁻/⁻ and Spry1⁻/⁻; Spry2⁻/⁻ mutant embryos by 17 - 20 s, and the size of the otic placode is restored to normal. Preliminary data suggest that a reduction in cell proliferation may help to reduce the size of an enlarged otic placode. Experiments exploring other mechanisms that explain the size correction of the otic placode are ongoing and will be presented.

Conclusion

These data suggest that organ-intrinsic mechanisms, such as modulation of cell proliferation, exist to correct aberrant enlargements of the otic placode, even after these cells have committed to an otic fate.

Retinoic Acid Signaling Regulates Tonotopic Patterning in the Developing Chick Cochlea

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Background

Frequency tuning depends on the basilar membrane’s resonance properties and the responses of hair cells (HCs). In the chick cochlea, there is a longitudinal gradient of HC phenotypes. Proximal-end HCs contain 250 or more stereocilia that reach a maximum length of 1.5 um, while distal-end HCs contain 50 or fewer stereocilia that reach a maximum length of 5.5 um. At locations between those ends, the number and length of stereocilia grade from HC to HC. We sought to discover what signals pattern the development of this gradient of HC phenotypes.

Methods

Using transcriptome sequencing we sought to identify components of signaling pathways that are differentially expressed along the tonotopic axis of the E6.5 chick cochlea. We used qPCR to check these results and to evaluate expression at other ages.

To test whether treatments would affect phenotypes that are linked to HC position, we cultured cochleae from E6.5 embryos for 6 days in a control medium or media supplemented either with all-trans retinoic acid (RA) or citral, a RA synthesis inhibitor. We used Scanning EM to assess stereocilia length and number, qPCR to assess expression of the B1 subunit of the BK channel, and immunostaining to assess Calbindin expression (phenotypes that all differ along the longitudinal axis).

Results

In our analysis of the sequence data we discovered opposing longitudinal gradients of mRNA expression for an enzyme involved in the synthesis of RA and an enzyme involved in RA degradation. RALDH3, a RA synthesis enzyme, is
expressed in a proximal-to-distal descending gradient, while CYP26C1, a RA metabolizing enzyme, is expressed in a distal-to-proximal gradient. However, a reverse (distal-to-proximal) gradient of RALDH3 is expressed at E10 and persists in the P14 cochlea. Through immunostaining, we confirmed that RALDH3 protein is expressed in the sensory epithelium.

When we cultured E6.5 cochleae in control medium for 6 days they developed the expected longitudinal gradient of HC phenotypes. In contrast, cochleae we cultured with RA developed HCs with distal-like phenotypes throughout the sensory epithelium. Cochleae cultured with the RA synthesis inhibitor, citral, developed HCs with more proximal-like phenotypes.

Conclusion
RA is a candidate morphogen that may influence the longitudinal patterning of HC phenotypes in the developing chicken cochlea. The evidence for a RA gradient we observed suggests that it may have an important role in the development of tonotopy.

A gradient of Bmp7 regulates tonotopic development in the chick basilar papilla
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Background
The auditory system of vertebrates that perceive sound in air is organized based on the separation of sounds into their component frequencies. Tonotopy, is present in the sensory epithelia of the mammalian cochlea and avian/reptilian basilar papilla (BP), as well as in higher auditory structures including the brain stem and cortex. Manifestations of tonotopy along the longitudinal axis of the cochlea/BP vary between classes with changes in basement membrane characteristics predominating in mammals, while gradients in hair cell phenotype are foremost in birds and reptiles. Despite the significance of tonotopic organization for auditory function, the molecular and cellular factors underlying development of tonotopic organization remain unknown.

Methods
To identify signaling molecules that regulate positional identity along the chick BP, we determined the developmental timing for acquisition of positional identity (E6.5) using known markers for tonotopy and then Affymetrix expression arrays to compare gene expression between proximal and distal halves of the BP. Amongst those genes differentially expressed, Bmp7, in particular, displayed a significant expression gradient from distal to proximal. Graded expression was confirmed using both in situ hybridization and quantitative real-time qPCR and was found to persist even in the post-hatch BP (P12). To determine the effects of disruption of the Bmp7 gradient, in vitro explant cultures were established at E6.5 and incubated for 6 days in either control media, or media supplemented with either Bmp7 or the Bmp antagonist, Noggin. Changes in positional identity along the tonotopic axis were assessed using known gradients in hair cell morphology including hair cell density, stereocilia length and number and the expression levels and cellular localization of Calbindin. Potential downstream targets of Bmp7 were identified by western blot and qPCR.

Results
Contrast to control explants, which developed a normal tonotopic gradient, explants maintained in media with a uniform concentration of Bmp7 showed changes in tonotopic phenotypes consistent with a conversion of the entire BP to a distal identity. Similarly, treatment with Noggin converted the entire BP to a proximal phenotype. Furthermore, analysis of downstream targets suggests that Bmp7 acts through interactions with the FGF and MAPK signalling pathways.

Conclusion
Bmp7 is expressed in a gradient along the tonotopic axis of the chick BP and disruption of that gradient induces phenotypic changes consistent with alterations in positional identity. These results suggest that Bmp7 signaling plays a major role in specification of hair cell positional identity, and tonotopic organization, along the BP.

ephrin-B2 is Required for Proper Development of an Inner Ear Transport Epithelium
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Background
Identification of shaker/waltzer behavior and a collapsed labyrinth in mice lacking EphB2, as well as in CD1 mice heterozygous for deletion of the ephrin-B2 (Efnb2) C-terminus, suggested a role for B-class Eph-ephrin signaling in
endolymph homeostasis. This was confirmed by the finding of decreased utricular [K+] in circling Efnb2 C-terminal deletion heterozygotes. However, the effects of these perturbations on the developing ear’s transport epithelia remained undefined.

Methods
We have generated mice that lack Efnb2 in developing ear tissues (Efnb2 CKO) and are viable until shortly after birth.

Results
At an early stage, Efnb2 CKO embryos display delayed outgrowth of the endolymphatic epithelium from the otocyst; this is preceded by a decreased rate of cell proliferation and heightened apoptosis. Delayed appearance of the structure is accompanied by disrupted patterning of endolymphatic duct and endolymphatic sac epithelia, as determined by mis-expression of genes in the Gbx, Dlx, and Wnt families, and tissue-level mis-localization of protein and mRNA elements in an ionocyte differentiation pathway and the Notch signaling pathway. In the CKO, mRNA and protein components of the ionocyte differentiation and Notch signaling pathways are altered with respect to cellular-level abundance, but sub-cellular localization of ion transport proteins and transcription factors is maintained. Components of the dysregulated ionocyte differentiation pathway include: the anion exchanger Pendrin (Slc26a4), mutations of which account for an estimated 5-8% of hereditary pre-lingual deafness in humans; the B1 subunit of an ATP-dependent proton pump (Atp6v1b1), mutations of which cause Renal Tubular Acidosis with Hearing Loss in humans; and Foxi1, a transcription factor and only known activator of Slc26a4 and Atp6v1b1 transcription in the ear. The utricle and saccule of Efnb2 CKO fetuses display giant otocoia and abnormal aggregations of pre-otoconial matter, which we interpret as morphological markers of endolymph abnormality. We also investigated the relevance of these fetal CKO phenotypes to balance disturbance in adult mice. Tissue-level mis-localization of the ionocyte differentiation pathway was deemed a quantitative trait, as its metrics are significantly increased in Efnb2 C-terminus deletion heterozygous fetuses of the CD1 circling strain compared with Efnb2 C-del heterozygous fetuses of a non-circling strain.

Conclusion
These analyses, together with characterizations of Efnb2 protein and mRNA expression, indicate that Efnb2 is required for proper growth, differentiation, and activity of the endolymphatic epithelium over a wide period of embryonic development.
Secreted Semaphorins Control SGN Peripheral Axon Motility and Hair Cell Innervation

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Background
During cochlear development, peripheral axons from spiral ganglion neurons (SGNs) navigate through different cell types before synapsing with inner or outer hair cells (IHCs or OHCs). Projections from type I SGNs (95% of the total population) form synapses with IHCs, whereas projections from type II SGNs (the remaining 5%) extend past IHCs and pillar cells before innervating OHCs. In addition, after extending into the OHC layer, type II processes turn abruptly toward the cochlear base. The guidance mechanisms that mediate these projection patterns are poorly understood. In these studies, we have found that class 3 secreted Semaphorins (Sema3s), which bind Neuropilin/Plexin (Nrp/Plxn) co-receptor complexes, may play roles in type I and type II SGN navigation.

Methods
To determine expression patterns for these factors, we used both in situ hybridization and immunostaining. To investigate the function of Sema3-Nrp/Plxn interactions in the cochlea, a combination of in vitro culture assays and mutant mouse models were examined. In addition, specific changes in embryonic SGN growth cone behaviors in response modulating Sema3-Nrp/Plxn interactions were visualized using a live imaging mouse model in which sparse numbers of SGNs express a red fluorophore (NgnCreERT2; R26R-tdTomato) while all hair cells express a green fluorophore (Atoh1-nGFP).

Results
Developing SGNs express Nrp1, Nrp2 and PlxnA3 whereas specific cell types within the cochlear epithelium express several Sema3s. Sema3a is expressed by cells of the lesser epithelial ridge and appears in an apex-to-base concentration gradient. Sema3f is expressed by lateral support cells within the organ of Corti. The loss of Semaphorin-Nrp signaling in Nrp mutant mouse models or by Nrp function-blocking antibodies in vitro leads to increased numbers of SGN peripheral processes in the OHC region and abundant type II SGNs that do not turn toward the base. Using our imaging model, we determined that adding exogenous Sema3A or Sema3F leads to the collapse of different subsets of SGN growth cones suggesting distinct roles for these guidance cues during cochlear development.

Conclusion
Overall, these data suggest that secreted Semaphorin-mediated activation of Nrp/Plxn complexes may act to restrict type I SGN processes to the IHC region. These data also reveal a signaling system that drives the apex-to-base migration of type II spiraling fibers.
Generation of ‘Three-Eared’ Frogs Reveals Molecular and Activity-Based Guidance of Central Projections
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Background
As new sensory organs arose in evolution, such as the ear, connections between the sensory system and CNS were formed. How these inner ear neurons find their central target in the hindbrain to transmit sound and movement information is not yet known. Development of the inner ear hair cells and neurons depends upon transcription factors such as Pax2/5/8 and several basic helix-loop-helix (bHLH) atonal family members. In the well-studied visual system, retinal ganglion cells (RGCs) similarly depend upon Pax2 and a bHLH atonal family member for their development. Given the molecular similarities of inner ear and eye development and the ability of inner ear afferents and RGCs to form a space-map in the CNS, it seems likely that the ear and eye use similar mechanisms for axon guidance. Projections of RGCs to the superior colliculus/tectum are guided by molecular cues (Eph/Ephrin) as well as activity-based mechanisms (formation of ocular dominance columns) in order to produce a topographical map of the environment. In this study, we generated ‘three-eared’ frogs in order to determine the amount of overlap or segregation of axonal projections when two ears are forced to innervate the same area of the hindbrain.

Methods
We transplanted an additional ear rostral to the native ear, either maintaining its orientation or rotating by 90 degrees. This allowed us to have the same or differential activity between the two ears, respectively. Swimming behavior was observed and axonal projections were labeled by implanting lipophilic dyes into the native and transplanted ears.

Results
Embryos in which the transplanted ear was in line with the native ear swam normally, whereas embryos in which the transplanted ear was rotated by 90 degrees swam aberrantly. Afferent axons from the two ears projected to overlapping areas when the transplanted ear was in line with the native ear. In contrast, when the transplanted ear was rotated by 90 degrees, afferent axons from the two ears were segregated from each other, forming ‘vestibular dominance columns’ reminiscent of the ocular dominance columns formed from varying activity between the two eyes.

Conclusion
These results suggest that afferents of transplanted ears rotated 90 degrees interfere with vestibular processing as indicated by the aberrant swimming. In addition, the partial overlap and segregation of afferent innervation in the two ‘three-eared’ frog models implies that both molecular and activity-based mechanisms affect central projections.

Identification of Multiple Marshalin Isoforms and Their Expression During Cochlear Development
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Background
The organ of Corti is composed of a group of highly polarized hair cells and their surrounding supporting cells, the latter having an unusual cytoskeletal structure. Microtubules (MT) and their networks play crucial roles during the development of the organ of Corti, which derives from simple epithelial cells. Marshalin has been identified as a MT minus-end-binding protein. MTs are present in loose dynamic networks in HCs (Slepecky et al., 1995), and tightly packed stable bundles in SCs (Angelborg and Engstrom, 1972). Although it is not known what factors contribute to form these distinctive patterns, we propose that Marshalin may be involved.

Methods
RNA was isolated using an Absolutely RNA®RT-PCR Miniprep Kit (Stratagene). RNA quality was measured by a 2100 Bioanalyzer (Agilent). Reverse transcription was performed by thermostable reversed transcriptase (Roche) at 55oC for one hour followed by PCR reactions using various primer sets. After cloning Marshalin cDNA into a mammalian expressing vector pEGFP-N2, plasmids encoding various marshalin isoforms were transiently transfected into opossum kidney (OK) cells. Marshalin-induced cytoskeletal changes were investigated using immunofluorescence.

Results
We discovered eight marshalin isoforms ranging from 863 to 1280 amino acids and carrying various protein-protein interacting domains including coiled-coil (CC), calponin homology, proline-rich (PR), and MT-binding domains referred to
as CKK. We examined structural changes in the cytoskeleton induced by expressing two of these marshalin isoforms in OK cells. Marshalin containing CC and PR domains induce thick structures with marshalin surrounding MT-containing bundles. Marshalin isoforms lacking these CC and PR domains also induce MT-based bundles but in a more slender form. These data suggest that marshalin is an important scaffolding protein, capable of modifying cytoskeletal networks through its interaction with various protein partners. To further investigate the function of different isoforms in vivo, we examined marshalin mRNA expression during organ of Corti development. Marshalin mRNA associated with each of the eight isoforms is detected at different developmental stages ranging from E17 to adult.

**Conclusion**

These data suggest that marshalin isoforms are abundantly expressed in the mammalian cochlea and that they play various roles in regulating cytoskeletal structures. The expression of one or more specific domains allows marshalin to interact with different protein partners to shape and maintain the architecture of the organ of Corti. (Work supported by NIH Grants DC011813, DC010633, DC00089, and the Knowles Hearing Center).

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**MafB is necessary for hearing function and auditory synapse development**

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**Background**

Auditory information is transmitted from the inner ear to the brain by spiral ganglion neurons (SGNs). To faithfully transmit sound information from hair cells to the brain, developing SGNs form a specialized synapse on hair cells, the ribbon synapse. The ribbon synapse is specialized for encoding acoustic signals with high temporal precision over a long period of time. Studying the molecular mechanism underlying how ribbon synapses develop may help us understand how sound information is transmitted and may lead to the development of new therapies for patients with hearing loss. We have identified the transcription factor MafB as a potential master regulator of auditory synapse development based on its specific expression in SGNs, functions in other developing systems, and ability to control the expression of known synaptic molecules.

**Methods**

To investigate the function of MafB in auditory synapse development, we generated a floxed allele of MafB and specifically removed MafB protein from SGNs. To complement MafB conditional knock-outs (MafBCKO), we created a strain of mice (MafBOE) that prematurely and excessively express MafB upon Cre-mediated recombination.

**Results**

MafBCKO mice are viable and exhibit no obvious behavioral abnormalities. Analysis of auditory function in MafBCKO mice by auditory brainstem responses shows that the mutants can still detect sound even at low intensities, consistent with the normal preservation of hair cell function seen upon measuring DPOAEs. However, the neural response was significantly decreased and delayed relative to controls, suggesting that MafB is required for proper SGN activity. Immunostaining for the presynaptic ribbon marker RIBEYE/CtBP2 and postsynaptic AMPA receptor subunit GluR2 revealed that the number of ribbon synapses is reduced in MafBCKO animals. In contrast to MafBCKO, MafBOE show increased ribbon synapses with altered cellular distribution. MafBCKO mice show normal SGN firing properties and have an intact olivocochlear efferent system, indicating that the reduction of ribbon synapses is not a consequence of impaired activity of SGNs or the efferent system.

**Conclusion**

These results suggest that MafB is required for normal formation of synapses between hair cells and SGNs. More detailed analyses of the MafB mutant mouse models together with genome wide studies to identify MafB downstream genes will be performed to help us understand how MafB contributes to the development of auditory synapses.
An overview of IGF signaling
Derek LeRoith, Derek LeRoith
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The insulin/insulin-like growth (IGF) system comprises insulin, IGF-1 and dIGF-2 as the ligands, six IGF binding proteins (IGFBPs) and three receptors; two receptors are functional as tyrosine kinases, the IR and the IGF-1R. Most of the functional activity of the IGF ligands occurs via the IGF-1R, a transmembrane tyrosine kinase receptor. Activation of the IGF-1R leads to activation of the PI3Kinase and MAPKiasse pathways that are important for growth and development, cell survival and certain differentiate functions in adult tissues. The IGF system is ubiquitous affecting almost every function of the body. Numerous gene-deletion mouse models of the IGFs, the IGFBPs and the IGF-1R have been created to established the whole body and tissue-specific effects of the IGF system, including the nervous system. In this presentation, we will address the major signaling processes involved in cellular function and the effects of deletion or overexpression of the IGF system in tissue function focusing where applicable on the nervous system.

Regulation of olfactory stem cell self-renewal and differentiation
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Tissue regeneration is a complex process that requires the coordination of stem cell proliferation and differentiation to maintain or repair the structure. The olfactory epithelium is a sensory neuroepithelium whose constituent cell types – including the olfactory sensory neurons – are continuously replaced during the lifetime of the animal. Following severe injury that results in the loss of mature cell types, the olfactory epithelium is reconstituted by the proliferation and differentiation of adult tissue stem cells. The regenerative capacity and limited number of cell types make the olfactory epithelium an excellent model for investigating stem cell regulation in vivo. Previous studies have identified the horizontal basal cell (HBC) as the multipotent neural stem cell of the olfactory epithelium; the molecules and pathways regulating this adult tissue stem cell are unknown, however. Using whole genome expression profiling of FACS-purified HBCs, we characterized the mRNA and miRNA transcriptomes of HBCs under conditions of quiescence and proliferation/differentiation. Through these studies we identified groups of genes associated with different phases of the HBC life cycle. In addition, we found that p63, a member of the p53 tumor suppressor gene family, is highly enriched in quiescent HBCs. p63 is a key regulator of stem cell self-renewal and differentiation in all stratified epithelia investigated to date. Through conditional inactivation of the p63 gene in HBCs we found that p63 is required cell-autonomously for olfactory stem cell renewal and functions to repress HBC differentiation. These results demonstrate a critical role of p63 in olfactory stem cell renewal and differentiation, where p63 serves as a "molecular switch" that controls these two alternate cell fates. Our studies provide the first molecular insights into the genetic network regulating stem cell dynamics in the olfactory epithelium and reveal an unexpected parallel between stem cell regulation in this sensory neuroepithelium and other epithelial tissues. Current work is focusing on identifying the downstream targets and interaction partners of p63 in the regulation of olfactory stem cell dynamics.
IGF signaling regulates cellular differentiation in the organ of Corti
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Insulin-like growth factor signaling is known to regulate multiple aspects of embryonic development including proliferation, growth, and differentiation. Mutations in members of the Igf signaling pathway lead to deafness in both humans and mice, demonstrating a role for this pathway in auditory function. While deletion of Igf1 results in post-natal loss of spiral ganglion neurons, any other effects of Igf signaling within the developing sensory epithelium were not known.

In situ hybridization for members of the Igf signaling pathway during embryonic cochlear development indicated complex patterns of expression for Igf1 and 2, Igf1r, and six Igf binding proteins. Igf1 and 2 are strongly expressed in Kölliker’s organ while Igf1r is expressed within the developing cochlear sensory epithelium.

Deletion of Igf1r results in shortened cochleae and defects in formation of the lateral and posterior semi-circular canals. Within the organ of Corti, Igf1r mutants exhibit decreased numbers of hair cells and a delay in hair cell differentiation. Analysis of the expression of developmental marker genes, including Sox2, p27kip1, and Atoh1 suggests that prosensory development is largely unaffected in Igf1r mutants, but that the transition from prosensory cell to hair cell is significantly delayed. Further analysis of Igf signaling in cochlear explants indicates independent effects on both cochlear elongation and hair cell differentiation, and suggests that the effects of Igf in the cochlea are regulated through PI3 kinase/AKT signaling.

Igf binding proteins can act as either enhancers or antagonists of Igf signaling. Igfbp3 expression is restricted to the prosensory domain prior to E14, but becomes down regulated in cells that develop as hair cells. Disruption of Notch signaling results in a loss of Igfbp3 as well as early differentiation of hair cells. These results imply that Igfbp3 could play a role in the timing of hair cell differentiation, possibly through antagonism of Igf signaling within the ear.

Overall, these results demonstrate an important role for Igf signaling in regulation of the transition between uncommitted prosensory cell and differentiating hair cell. Defects arising from disruption of this pathway could underlie the hearing loss observed in both humans and mice with perturbations of Igf signaling.

Insulin-like growth factor 1 deficiency and hearing loss
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Insulin like growth factor 1 (IGF-1) plays a central role in embryonic development and adult tissue homeostasis. Accordingly, it is an essential factor for the regulation of cochlear development through controlling apoptosis and late neuronal differentiation. In the cochlea, IGF-1 actions are mediated by a network of protein kinases (RAF, AKT and p38 MAPK) that modulate the expression and activity of transcription factors (AP1, MEF2, FoxG1 and FoxM1) leading to the regulation of cell cycle and metabolism.

IGF-1 deficiency causes hearing loss in mice and men, whereas low serum levels of IGF-1 are associated with human syndromes showing hearing loss and with premature presbyacusis. Animal models are fundamental to understand the genetic, epigenetic, and environmental factors that contribute to human hearing loss. Mouse IGF-1 plasmatic levels decrease with ageing, concomitantly hearing loss and retinal degeneration occur, in a normal ageing process that is accelerated in the heterozygous Igf1⁺/⁻ mouse. Furthermore, low IGF-1 levels predispose to increased damage after noise-exposure. In the Igf1 null mouse, hearing-loss is caused by early neuronal loss and age-related stria vascularis alterations.

In summary, our data suggest that serum IGF-1 levels could be a novel diagnostic tool for hearing loss and support the hypothesis that IGF-1-based treatments have potential for the protection or repair of hearing loss.
Clinical efficacy of topical IGF1 treatment for sudden deafness

Takayuki Nakagawa

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Background

Several works from different laboratories have indicated that the insulin-like growth factor (IGF) signaling plays important roles in embryonic development and adult homeostasis of the cochlea. These findings encouraged us to investigate the efficacy of IGF1 in the treatment of sudden sensorineural hearing loss (SSHL). For this purpose, we have developed the drug delivery system using gelatin hydrogels for sustained, topical delivery of IGF1 into the cochlear fluid. Here we report clinical efficacy of topical IGF1 application via gelatin hydrogels for SSHL patients refractory to systemic steroid application.

Methods

A prospective, single-armed trial was designed using a historical control of hyperbaric oxygen therapy. Patients with SSHL that showed no recovery to systemic glucocorticoid administration were recruited. We applied gelatin hydrogels impregnated with recombinant human IGF1 in the round window niche. The primary outcome measure was the proportion of patients showing hearing improvement 12 weeks after the test treatment. The secondary outcome measures were the proportion of patients showing improvement at 24 weeks and the incidence of adverse events. We also performed retrospective chart review of these patients to figure out alterations in thresholds of pure tone audiometry (PTA).

Results

In total, 25 patients received the test treatment at a median of 23 days after the onset of SSHL. At 12 weeks after the test treatment, 48% (P = 0.086) of patients showed hearing improvement, and the proportion increased to 56% (P = 0.015) at 24 weeks. No serious adverse events were observed. These findings indicate that topical IGF1 application using gelatin hydrogels is well tolerated and may be efficacious for hearing recovery in patients with SSHL refractory to systemic steroids. In analyses of audiometric alterations, significant improvements of PTA thresholds were identified. Interestingly, the time course of hearing recovery is very slow. The major recovery of hearing appeared 4 weeks after application, and a few patients demonstrated further recovery of hearing until 24 weeks after treatment. These findings suggest that not only protection of hair cells but also other regenerative processes may be involved in mechanisms for hearing recovery.

Conclusions

A clinical trial of topical IGF1 application in SSHL patients indicates the potential of IGF1 as a therapeutic agent for SSHL, and the need of further basic experiments to explore mechanisms for hearing recovery by IGF1.
Mechanisms of cochlear hair cell protection by IGF-1
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We have investigated the role of insulin like growth factor-1 (IGF-1) in cochlear hair cell protection. IGF-1 protects mammalian cochlear hair cells from noise exposure or from ischemic stress. The clinical trial for the treatment of idiopathic sudden sensorineural hearing loss cases that was refractory to the systemic steroid treatment showed improved hearing threshold in 56% of patients who were treated with IGF-1.

To establish more effective protocols of IGF-1 therapy for sensorineural hearing loss we tried to elucidate the mechanisms of cochlear hair cell protection by IGF-1 using cochlear explant culture of neonatal mice. We used neomycin as a method of hair cell impairment that was protected by IGF-1 in the explant culture system.

This system enables us to add various inhibitors at the most effective timing and concentration to clarify the downstream signal pathways of IGF-1 exerting the cochlear hair cell protection. We examined inhibitors of PI3K, Akt, MEK, and PKC that constitute the components of major downstream signal pathway of IGF-1. All tested inhibitors attenuated the effect of IGF-1 although the Akt inhibitor only attenuated the protection of inner hair cells, indicating that IGF-1 treatment could activate several kinds of downstream signal pathways. The specific action of Akt was confirmed by immunohistochemistry of phospho-Akt that was detected around inner hair cells. To clarify the effector of IGF-1 in the protection of hair cells, we performed comprehensive gene expression profiling using microarray. We identified two specific effector genes of IGF-1 signal, Gap-43 and N tissue1, in our experimental system.

To elucidate the cellular mechanisms involved in IGF-1 action, we examined the apoptosis and cell proliferation status after IGF-1 treatment. Decrease of apoptotic hair cell numbers and proliferation of Hensen’s cells and Claudius’ cells were observed when explant culture impaired by neomycin was treated with IGF-1. Contribution of the supporting cell proliferation to the maintenance of outer hair cells was confirmed by treatment with two different proliferation inhibitors, aphidicholin and mimosine.

In conclusion, these results indicated that IGF-1 activated several downstream cascades and, as a result, it promoted the proliferation of supporting cells and inhibited the apoptosis of cochlear hair cells in neonatal mice.
Identification of novel downstream effectors of IGF-1 signal pathways using a comprehensive gene expression analysis

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Background
We have revealed that insulin-like growth factor1 (IGF-1) protects cochlear hair cells of neonatal mice against aminoglycoside via both the phosphatidylinositol 3-kinase (PI3K)/Akt pathway and the mitogen-activated protein kinase kinase /extracellular signal-regulated kinase (MEK/ERK) pathway. However the effector genes of PI3K/Akt and MEK/ERK which work directly on cochlear hair cell protection have not been unveiled so far. To identify the effectors of IGF-1, we screened whole sets of expressed genes in the cochlear explant cultures using microarray and confirmed the results using quantitative RT-PCR (qRT-PCR).

Methods
1. Total RNA was extracted from the cochlear explant cultures of neonatal mice that were treated with only neomycin or both neomycin and IGF-1 for various durations. cDNAs were prepared from the total RNA and the expression levels of whole genes were compared between the control and IGF-1-treated groups using microarray provided by Affymetrix. We picked up the genes whose expression levels increased twofold in the experimental groups compared with those in the control groups. The differences in the expression levels were confirmed by qRT-PCR.
2. To confirm the identified genes were really at the downstream of the IGF-1 pathways, the changes of their expression levels after treatment with AKT or MEK inhibitor were tested using qRT-PCR.

Results
The expression levels of Gap43 and Ntn1 were up-regulated in the IGF-1-treated cochlear explant cultures compared with the control samples. The expression levels of Gap43 and Ntn1 were significantly reduced by the addition of Akt inhibitor or MEK inhibitor.

Conclusion
As effectors of IGF-1 signaling in the context of hair cell protection from aminoglycoside, we identified two kinds of genes, Gap43 and Ntn1 whose expression level increase was canceled by the addition of the PI3K or MEK inhibitor. These findings indicated that Gap43 and Ntn1 are the downstream effectors of IGF-1 in the cochlear sensory epithelium that are under the regulation of both the PI3K/Akt and MEK/ERK pathways.

Lmo4 signaling in cisplatin mediated ototoxicity

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Background
The transcriptional regulator Lmo4 was recently identified as the major nitrated cochlear protein in cisplatin ototoxicity. Nitrated Lmo4 was localized in a number of critical cell types including outer hair cells, spiral ganglion neurons, and striae cells. Moreover, cisplatin treatment significantly decreased Lmo4 levels in the cochlea. The functional significance of a decrease in Lmo4 levels was indicated by an increased expression of activated caspase 3 in the inner and outer hair cells, upon silencing cochlear Lmo4 with siRNAs. These findings demonstrated the crucial role of Lmo4 in mediating cisplatin ototoxicity. However, the molecular signaling associated with cisplatin-induced modulation of cochlear Lmo4 is poorly understood.

Methods
Wistar rats were treated with a single dose of 16 mg/kg cisplatin and cochlear expression of mRNAs and proteins associated with Lmo4 signaling were evaluated, 3 days post treatment. A custom gene array was designed to investigate at least ten known binding partners and downstream targets of Lmo4, while proteins of interest were assessed by immunoblots. Co-treatment with 100 mg/kg/day Trolox was used to inhibit cochlear nitrosative stress and prevent cisplatin-induced ototoxicity.

Results
RT-PCR analysis of cochlear mRNA levels, indicated that cisplatin treatment modulated the expression levels of several binding partners and targets of Lmo4. Particularly, the cochlear expression of Esr1 was significantly up-regulated by cisplatin treatment, while the expression of Jak1 and Stat3 was down-regulated. Co-treatment with antioxidant Trolox, which is also an inhibitor of peroxynitrite, attenuated their cisplatin-induced modulation in the cochlea. Further evaluation
of the cochlear protein levels of Stat3, a downstream target of Lmo4 that is implicated in drug mediated apoptosis, supported the observed changes in the mRNA levels. Cisplatin treatment significantly decreased Stat3 protein expression (p<0.001), while co-treatment with Trolox attenuated the cisplatin-induced decrease in Stat3 expression, in the cochlea (p<0.01). Trolox co-treatment has previously been reported to prevent cisplatin-induced hearing loss. Therefore, these results illuminate a potential role of the key downstream players, Jak1 and Stat3, in cochlear Lmo4 signaling and implicate their importance in mediating the ototoxic effects of cisplatin.

**Conclusion**

Based on these findings we hypothesize that cisplatin-induced nitration of Lmo4 decreases the cochlear expression of Lmo4 and facilitates the modulation of its binding partners Esr1 and Jak1, which in turn represses Stat3 and augments cochlear apoptosis. (Funded by NIH/NIDCD: R03DC010225, SJ)

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**Oral Capsaicin Prevents Cisplatin Ototoxicity**

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**Background**

Ototoxicity and nephrotoxicity are dose-limiting side effects of cisplatin, a widely used anticancer drug. Data from our laboratory suggest that cisplatin ototoxicity is linked, in part, to activation of transient receptor potential vanilloid 1 (TRPV1) receptor in the cochlea. Direct activation of TRPV1 by capsaicin (a TRPV1 agonist) produced transient hearing loss (~25 dB shifts) for 24h, which recovered by 72 h. This temporary threshold shift was not associated with damage or loss of outer hair cells. We found increases in stress response genes like TRPV1, NOX3 NADPH oxidase and pro-inflammatory moieties like iNOS, COX2 and TNF-α, but no increases in pro-apoptotic molecules like p53 or bax.

Furthermore, capsaicin administration by the trans-tympanic route (50µl of 0.1µM) 24h prior to cisplatin (11 mg/kg, i.p) attenuated cisplatin induced hearing loss. We therefore hypothesized that oral administration of capsaicin would also help ameliorate cisplatin induced hearing loss.

**Methods**

Male Wistar rats were used for this study. Baseline ABRs were recorded, capsaicin or PBS was administered by oral gavage (2% in water, 1ml) 24h prior to cisplatin administration (11 mg/kg, i.p). Post treatment ABRs were recorded 72 h after cisplatin administration. Cochleae were collected for morphological analysis by scanning electron microscopy (SEM), gene expression by real time RT-PCR and immunohistochemistry of the mid-modiolar sections of the rat cochleae.

**Results**

Cisplatin treatment produced ~25 dB shift in hearing threshold. This cisplatin induced hearing loss was attenuated to (<10 dB) by both trans-tympanic as well as oral capsaicin pre-treatment. Preservation of hearing was associated with no significant damage or loss of OHCs in rats pretreated with trans-tympanic capsaicin prior to cisplatin administration as seen morphologically in SEM images, when compared to rats treated with vehicle plus cisplatin.

**Conclusion**

Oral capsaicin (levels equivalent to those when consuming spicy food) can attenuate cisplatin induced hearing loss. To our knowledge this is the first report showing the efficacy of oral capsaicin works as a pre-conditioning stimulus that ameliorates cisplatin ototoxicity. (Supported by NIH grants R01DC02396 to LPR, funds from SIU School of Medicine).

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**Effect of somatostatin and octreotide on spiral ganglion neurites and gentamicin-induced auditory hair cell loss**

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**Background**

Somatostatin (SST), a peptide with hormone/neurotransmitter properties is known to affect targets in the CNS. It has been shown that SST receptors are expressed in the developing spiral ganglion and organ of Corti (OC) as well in the adult mammalian inner ear. SST has previously been shown to protect against gentamicin-induced auditory hair cell loss in vitro. We hypothesize that SST plays a role in spiral ganglion (SG) neurite outgrowth and that octreotide, a clinically used somatostatin analog, can protect hair cells from gentamicin-induced damage. Therefore the effect of SST and octreotide
on SG neurite length and number was analyzed. In addition, we assessed the effect of octreotide on gentamicin-induced hair cell loss in vitro.

**Methods**
OC and SG explants were extracted from 5-day old wistar rats. SG explants were treated with different concentrations of SST or octreotide. Protection of auditory hair cells from gentamicin-induced toxicity was tested using two concentrations of octreotide. OC explants were preincubated with or without octreotide following by addition of gentamicine.

**Results**
Octreotide and SST increased SG neuronal survival. We observed more neurites per spiral ganglion explants treated in the SST and octreotide samples compared to control. The effect was dose dependent and statistical significant (ANOVA, p<.05). The length of neurites was slightly decreased at the lowest SST concentration (1uM) and at all octreotide concentrations (1uM-20uM) used compared to controls (ANOVA, p<.05). There was significantly less hair cell loss in OC treated with gentamicin in presence of octreotide compared to samples treated with gentamicin alone (ANOVA, p<.01).

**Conclusion**
Decreased hair cell loss in octreotide-treated samples that had been exposed to gentamicin provides evidence for a protective effect of octreotide in aminoglycoside-induced hair cell death in vitro. The effect of SST and octreotide on both length and number of SG neurites in vitro points to a possible involvement of SST in cochlear development.

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**308 Role of STAT1-p53 Axis in Mediating Gentamicin-Induced Cellular Stress and Hair Cell Death**

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**Background**
The aminoglycoside antibiotics, including the prototype gentamicin, are a well known cause of irreversible hearing loss. Reactive oxygen species (ROS) formation within the cochlea is a critical event for ototoxicity and hair cell death. In cisplatin-treated rat cochleae, ROS lead to activation of signal transducer and activator of transcription 1 (STAT1) and p53-mediated killing. We therefore hypothesized that ROS activation of the STAT1 – p53 axis may also mediate hair cell death after gentamicin therapy. We further predicted that pretreatment with the putative otoprotectant transplatin could decrease cellular stress and prevent cell death.

**Methods**
All assays were conducted using UB/OC-1 cells, a conditionally immortalized cell line derived from the organ of Corti. Gentamicin was administered at 0, 100 µM, 500µM, or 1mM doses, and transplatin was administered at 0, 5, and 10 µM doses. Gene expression was determined by real-time RT-PCR. MTT Tetrazolium dye reduction proliferation assays were used for assessment of cell viability. ROS formation was evaluated with H2DCFDA dye fluorescence assay using confocal microscopy. For immunocytochemistry (ICC), UB/OC-1 cells were treated with gentamicin for 24 hours and probed for TNF-alpha, NADPH oxidase isoform-3 (NOX3), transient receptor potential vanilloid 1 (TRPV1), and p-STAT by immunofluorescence.

**Results**
Gentamicin treatment increased STAT1 and p53 expression on real time RT-PCR. MTT proliferation assay demonstrated a 15% decrease in cell viability with gentamicin versus untreated control cells at 48 hours. Gentamicin-induced ROS formation was confirmed by live imaging by fluorescence confocal microscopy, and NOX3 prevalence was also seen to be increased. Gentamicin also resulted in increased prevalence of phosphorylated STAT1 (pSTAT1) at Ser727 residue and pSTAT1-responsive genes on ICC. Translocation of pSTAT1 to cell nuclei was confirmed with merged images with nuclear stain (DAPI). The TRPV1 channel, thought to play a role in gentamicin uptake, was also upregulated by gentamicin. On MTT assay, pretreatment with transplatin prior to gentamicin conferred complete protection from cell death at 48hrs, with an absorbance profile equal to controls. Transplatin pretreatment also normalized prevalence of NOX3, pSTAT1, TRPV1, and TNF-alpha to baseline levels on ICC.

**Conclusion**
These data suggest that the STAT1-p53 axis mediates gentamicin-induced hair cell death, with transplatin conferring otoprotection. Transplatin prevents reactive oxygen species formation and attenuates cellular stress, providing a potential mechanism for decreased STAT1-p53 activation.
The JNK signaling complex in hair cell damage
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Background
Hair cell (HC) damage involves multiple molecular substrates. Among these, signaling via the c-Jun N-terminal kinase (JNK) pathway is thought to play a central role in noise, aminoglycoside and age-related HC loss.

Methods
We evaluated signaling molecules upstream of JNK using proteomics and specific inhibitors, in a gentamicin (GM)-induced in vitro model of HC loss. We also used proteomics in an in vivo model of noise-induced hearing loss, by comparing cochlear proteins from mice exposed to brief noise at an intensity that induces only temporary threshold shift (TTS), versus exposure to an intensity that induces permanent threshold shift (PTS). To assess the role of different JNK isoforms we evaluated, in vitro, GM-induced HC damage in mice deficient in JNK1 or JNK2.

Results
Inhibitor data indicated that GM activates small GTPases including kRas, Rac and cdc42. Inhibitors also implicated mixed lineage kinases and JNK itself. Comparison of proteins from cochleas with TTS versus those with PTS identified rapid, noise-induced reductions in several small GTPases of the Ras and Rho families in PTS cochleas compared to their levels in TTS cochleas. This is consistent with high levels of activation leading to degradation. A cdc42/Rho activating protein, a protein downstream from RhoA, and other JNK-interacting proteins were also reduced. Preliminary in vitro data from mice deficient in JNK1 or JNK2 indicate that JNK1 plays a protective role while JNK2 mediates HC damage. This illustrates the complex nature of signaling related to HC death, in which survival and damage pathways compete to determine the fate of the cell.

Conclusion
Our results support the idea that JNK signaling, including activation of upstream Ras and Rho family GTPases, is involved in HC damage due to both drugs and noise. In addition, we identified new molecular candidates for roles in regulating this pathway within the HC, providing additional targets for potential pharmacological manipulation, and potentially identified a protective JNK isoform.

Chemotherapy for the Future: Novel Aminoglycosides Dissecting Antibacterial Activity and Ototoxicity
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Background
Ototoxicity is thought to be inherent to aminoglycosides, compromising the clinical use of these important antibacterials. While we have shown clinically successful pharmacological mitigation of aminoglycoside-induced hearing loss (New Engl. J. Med. 354:1856-1857, 2006), chemotherapy of the future would benefit from aminoglycosides effective against multidrug resistant bacteria but with no or little ototoxic potential.

Methods
Chemical synthesis; in-vitro translation assays; crystallography; site directed mutagenesis; antibacterial activity; reactive oxygen species; ototoxicity in murine organ culture and in guinea pig in vivo.

Results
Our mechanistic concept postulates a key role for the mitochondrial ribosome (mitoribosome) in aminoglycoside ototoxicity (PNAS 105:20888-20893, 2008). We have previously reported on the finding (PNAS 109:10984-10989, 2012) that apramycin, a structurally unique aminoglycoside in veterinary oral use for treatment of intestinal infections, shows low activity towards eukaryotic ribosomes, including hybrid ribosomes carrying the aminoglycoside-susceptibility A1555G allele. In murine cochlear explants, apramycin caused only little hair cell damage. In guinea pig in vivo, and in contrast to gentamicin, ototoxicity developed more gradually and only at higher concentrations. In accordance with its lesser toxicity, apramycin did not elicit free-radicals (ROS) at low concentrations. However, for both gentamicin and apramycin, the ROS marker nitrotyrosine appeared at antibiotic levels that caused loss of hair cells. This correlation with ototoxicity – rather than with drug concentration – supports the notion that generation of ROS is an integral part of the mechanisms that lead
to cell death. Based on this proof-of-concept that antibacterial activity can be dissected from aminoglycoside ototoxicity, we have now developed aminoglycoside compounds with low ototoxic potential.

**Conclusion**

Together with three-dimensional structures of apramycin-ribosome complexes at 3.5 Å resolution, our results provide a conceptual framework to develop less toxic aminoglycosides by hypothesis-driven chemical synthesis. For the present, apramycin’s high efficacy against drug-resistant strains combined with low ototoxic potential makes it a drug of immediate clinical interest. New lead compounds promise a safety margin (antibacterial efficacy vs. ototoxicity) more than an order of magnitude better than gentamicin.

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**Cisplatin-induced influences on age-related changes in auditory physiology in the Fischer 344/NHsd rat**

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**Background**

Cisplatin (CDDP) was approved for use by the FDA in 1978. A large proportion of the oncology patients who underwent successful CDDP treatment did so early in their life spans. Therefore, there is a population of patients with CDDP-induced hearing loss, or who received CDDP without detectable threshold shift, who are now reaching their sixth, seventh, and eighth decades of life, the times when age-related hearing is expected to set on. This population prompted the current study in which Fischer 344/NHsd rats, a model of age-related hearing loss in the cochlea, were exposed to cisplatin at age 7 months. The quiet-aged Fischer 344/NHsd rat develops hearing loss in the high frequencies between the ages of 9 and 12 months of age, and the hearing loss spreads to mid and low frequencies as the animals age from 12 to 24 months.

**Methods**

Eight animals received a single dose of 7 mg/kg CDDP in saline delivered intraperitoneally (i.p.) in a 30-minute infusion, while six animals received i.p. saline only as controls. All animals were aged 7 months at the time of the CDDP exposure. Auditory physiology in the form of auditory brainstem response (ABR) thresholds and supra-threshold amplitude input-output functions, along with DPOAE input-output functions, were evaluated monthly to assess the development and magnitude of age-related hearing loss in the CDDP-exposed and saline control rats. Monthly testing was performed for 11 months after the exposures until the animals were 18 months of age.

**Results**

The ABR threshold results revealed an acute threshold shift at the highest frequencies tested (30-40 kHz) within one month after CDDP exposure, and then earlier onset and greater severity of age-related hearing loss in the mid- to high-frequency range (15-30 kHz) by age 15 months. ABR input-output functions also showed acute effects of CDDP, followed by gradual amplitude reductions over time in the CDDP-exposed ears.

**Conclusion**

The results suggest that CDDP ototoxicity early in adulthood alters the trajectory of age-related hearing loss in the Fischer 344/NHsd rat. The age-related hearing loss of the Fischer 344/NHsd rat is associated with outer hair cell pathology, but the accelerated age-related hearing loss observed in the current study cannot yet be linked to a specific underlying pathology. Anatomical studies of the cochleae from the subjects are ongoing.
Age-Related Changes in Inner Hair Cell – Auditory Nerve Connections and Ntf3 expression in the cochlea of UMHet4 mice

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Background
We found a significant decline in the number of inner hair cell – auditory nerve (IHC-AN) connections across turns by 22-24 months of age in normal female UMHet4 mice. There was no further decrease at 27-29 months of age. On the other hand, auditory brain stem response (ABR) threshold shifts and hair cell loss were largely confined to apical / low frequency regions at 22-24 months and both measures showed considerable further decrement at 27-29 months. This suggested differences in underlying mechanisms. Recent studies show a role for NT3/Ntf3 in inner hair cell – auditory nerve synapse development, maintenance, regulation and regeneration (Wang and Green, 2012, Wan et al, 2012). We therefore looked for changes in Ntf3 gene expression at 22-24 months as well as changes in Gdnf and Bdnf, which can serve as survival factors for spiral ganglion neurons.

Methods
UMHet4 mice are a genetically heterogeneous four-way cross between female MOLF/EiJ x 129S1/SvImJ F1 mice and male C3H/HeJ x FVB/NJ F1 mice, all are negative for the AhI1 allele that generates hearing loss appearing in young adults. Mice were tested at 5-7 months, 22-24 months and 27-29 months for ABR, assessed for hair cell loss along the cochlear spiral and for IHC-AN connections based on CTBP2 immunostaining. Gene expression of Ntf3, Bdnf and Gdnf in the cochlea was compared between 5-7 month and 22-24 month old animals using qRT-PCR.

Results
There was a significant decrease of well over 50% in Ntf3 (NT3) gene expression in the cochlea of 22-24 month old UMHet4 mice compared to 5-7 month mice. Bdnf showed a smaller decrease in gene expression with considerable variability such that it was not significant. There was no decrease in Gdnf gene expression.

Conclusion
Given the findings by Wan et al (2012) and Wang & Green (2012) that NT3 regulates and maintains IHC-AN connections, the large age-related decrease in Ntf3 gene expression we find in the mouse cochlea by 22-24 months of age could result in a decreased ability to maintain the full number of connections found in young animals. This could, in turn, explain our finding a 20-34% percent age-related decrease in IHC-AN connections.

Studies were supported by NIH grant AG025164 and NIDCD P30 grant DC005188
Hearing Loss and Changes in Brain Volume in Older Adults
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Background
Cross-sectional neuroimaging studies have demonstrated that audiometric hearing loss is associated with reduced volumes in the auditory cortex while epidemiologic studies have demonstrated that hearing loss is independently associated with accelerated cognitive decline and incident dementia. Whether hearing loss is associated with brain regions outside auditory cortex and with trajectories of brain volume decline is unknown.

Methods
We prospectively followed 126 individuals (ages 56-86) without prevalent dementia in the neuroimaging substudy of the Baltimore Longitudinal Study of Aging for a mean of 6.4 years with annual magnetic resonance brain scans. Audiometry was performed before the baseline scan, and hearing loss was defined as a speech-frequency pure tone average > 25 dB. Mixed-effects regression models were used to compare regional brain volume trajectories of individuals with hearing loss versus normal hearing.

Results
Accelerated volume declines over the follow-up period in whole brain, regional temporal lobe volumes (parahippocampus, superior, middle, and inferior temporal gyri), and cingulate gyrus were observed in individuals with hearing loss compared to those with normal hearing (p < .05). These results were robust to adjustment for multiple confounders and were consistent with voxel-based analyses.

Conclusion
Hearing loss is independently associated with accelerated whole brain and superior temporal gyrus atrophy among older adults. Whether hearing loss is an early marker or possibly a modifiable risk factor for structural brain aging deserves further study.

Cortical evoked potential measures as biomarkers of binaural temporal processing in older adults
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Background
Temporal envelope and fine structure cues in binaural tasks such as the masking level difference (MLD) provide measurable advantages for detecting signals in background noise. Compared to younger listeners, however, older adults often show reduced performance on such binaural tasks, thought to be due in part to diminished temporal processing with age and peripheral hearing loss. It is not clear to what extent diminished perceptual performance reflects age-related changes in central auditory encoding of envelope cues, fine structure cues or both. The purpose of this study was to evaluate the utility of cortical evoked potential measures as biomarkers of binaural temporal envelope and fine structure cues in an MLD paradigm in older adults.

Methods
Psychophysical and cortical evoked potential thresholds were measured for diotic (So) and dichotic (Sπ) tonal signals presented in continuous diotic (No) maskers in ten older adults with essentially flat, mild to moderate (35-55 dB HL) sensorineural hearing loss. The MLD was computed as the threshold difference in dichotic (NoSπ) versus diotic (NoSo) conditions. The contribution of stimulus envelope cues was examined using narrowband noise (50 Hz) with varying degrees of envelope fluctuation (Gaussian vs. Low-noise noise), and the contribution of fine structure cues was evaluated using low and high center frequencies (500 vs. 4000 Hz).

Results
Cortical evoked potential thresholds were within 10-15 dB of behavioral thresholds and highly correlated with behavioral thresholds. Both behavioral and evoked-potential MLDs were largest in the 500 Hz Gaussian noise condition where envelope and fine structure cues were available. MLDs were reduced in 500 Hz low-noise noise where envelope cues were minimized while fine structure cues were present. MLDs also were reduced in the 4000 Hz Gaussian noise condition where envelope cues were available but fine structure cues were not. Compared to young adults with normal hearing, MLDs in older adults were significantly smaller at 500 Hz in both Gaussian and low-noise noise, primarily due to
reduced masking release in dichotic conditions, but were minimally different for 4000 Hz conditions. Notably, N1 and P2 latencies were longer in Gaussian noise dichotic conditions for older compared to younger adults.

Conclusion
The results indicate that cortical evoked potential measures may have utility as biomarkers of binaural temporal processing in older adults and other clinical populations.

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**Genes on Chromosome 17 Contribute to Age-Related Hearing Loss in 129S6/SvEvTac**
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Background
Presbycusis is one of the main causes of acquired deafness. Mice with age-related hearing loss (AHL) are used as models of presbycusis because of their homology to the human auditory system. From the eight AHL loci described to date, three have led to the identification of genes that contribute to AHL. The 129S6/SvEvTac (S6) mouse is commonly used for gene targeting and noise resistance studies; however, S6 also has early-onset AHL. A similar AHL phenotype was mapped to Chromosome (Chr) 17 and Chr 10 in the genetically related 101H strain. Our objectives are: to use S6 to improve our understanding of AHL mechanisms, to identify the physical location of the AHL gene(s) for S6, and to determine whether 101H and S6 share similar genes that contribute to AHL.

Methods
Hearing experiments were conducted in S6 and 101H mice. We used auditory brainstem responses (ABRs) to measure hearing thresholds at six frequencies between 8khz and 40kHz. Single-nucleotide polymorphisms (SNPs) were used in combination with Taqman assay for genotyping. By selectively backcrossing S6 to a good hearing strain, CBA/CaJ (CB), we are creating six novel recombinant-inbred (congenic) CB mouse strains with fragments of Chr 17 from S6. We hypothesize that genes within Chr 17 contribute to AHL in the S6 strain.

Results
By 4 weeks of age, S6 mice have an average hearing loss of 22dB (SPL) between 24-40kHz compared to CB. A congenic strain with the proximal 30Mb of Chr 17 from S6, CB-S6-(prox-D17Mit29), has similar hearing loss of 23dB between 24-40kHz. The proximal 30Mb region of Chr 17 contains upwards of 500 known genes of which 9 have been implicated with hearing. New congenic lines were generated to localize the causal gene with higher resolution. Congenic mouse CB-S6-(rs1348286-rs13482938) and CB-S6-(rs13482886-rs13482938) maintain a high frequency hearing loss in the 24-40kHz region while CB-S6-(prox-rs13482886) has hearing threshold indistinguishable from CB.

Conclusion
These data locate the causal genes between 21Mb and 28Mb on Chr 17. A complementation test is being conducted between the above congenic lines to determine if 101H share a common allele with S6 that produces the AHL phenotype. These studies will help identify genes causing AHL thereby elucidating homologous genes in humans that may contribute to presbycusis.

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**The Aging Auditory System of the Common Marmoset (Callithrix jacchus)**
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Background
The purpose of this study was to determine the effects of aging on auditory function in common marmosets (Callithrix jacchus). This species has increased in popularity as a model for auditory research, but little is known about the effects of aging on auditory function in the marmoset.

Methods
Hearing sensitivity and neural function were assessed in 15 young (1-4 yrs), 10 older-adult (5-8 yrs), and 10 aged (8-12 yrs) males using auditory brainstem responses (ABRs). Cochlear function was assessed in the same subjects using
distortion-product otoacoustic emissions (DPOAEs). Significance of group differences was determined using effect size (Cohen's $d$) and bootstrap resampling.

ABR tone-burst stimuli were presented in 3-kHz steps from 3-9 and 15-24kHz at 74 dB SPL to assess the peak latency of waves I, II, and V. Tone-burst levels were reduced to determine threshold. Threshold was defined as the tone-burst level which elicited two repeatable waveforms (Pearson's $r > 0$, $p < 0.001$) with peaks that were distinguishable from the noise floor.

For DPOAEs, $f_2$ ranged from 3-24kHz in 1/5-octave steps ($f_2/f_1 = 1.21$), and $L_1$ was increased from 50-74 dB SPL in 6-dB steps.

**Results**

Consistent with the behavioral audiograms measured by other groups, ABR thresholds were lowest in each group at 6 kHz. ABR thresholds of older-adult males were significantly elevated at 3, 9, 15, and 18 kHz when compared to the young males. Thresholds of aged males were significantly elevated at all stimulus frequencies when compared to the young, and were significantly greater than older adults at 24kHz.

The ABR wave V latency, but not wave I and II latencies, was significantly prolonged in older-adults at 6, 18, and 21 kHz. In aged marmosets, wave I, II, and V latencies were delayed compared to the young males in a non-frequency specific manner.

DPOAE levels were reduced significantly in aged males, but not older-adult marmosets, at a wide range of frequencies and primary-tone levels.

**Conclusion**

These data suggest that the decline in hearing sensitivity observed in male marmosets does not progress from high- to low frequencies, that cochlear function may be preserved until late in life, and that central auditory dysfunction may precede peripheral deficits in this species.
Ventriloquism Effect in Young and Elderly Normal-Hearing Listeners

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\textbf{Background}

The success in fitting hearing aids depends, among others, on the ability to combine the auditory and visual information for speech perception. The audiovisual integration can be observed in the ventriloquist effect, where an observer localizes sound at a visually dominated spatial position. In this study, the suitability of the ventriloquist effect as a measure for the audiovisual integration was investigated. Assuming that elderly subjects have a greater ability for audiovisual integration, the effect of the age was also taken into account.

\textbf{Methods}

Minimum audible angles (MAAs) in the horizontal plane were measured in an acoustic lateralization task with a visual distracter. The acoustic stimulus was a 500-Hz pure tone presented via headphones at a lateral angle at the right or left side by filtering the stimulus with generic head-related transfer functions. Each lateral side was tested in an adaptive 3-up-1-down run, starting at the angle of 10\degree, and decreasing after three succeeding correct responses. Two runs, corresponding to the two sides, were randomly interleaved. The MAA was the difference between the lateral angles obtained from the two runs averaged over the last four reversals. The visual stimulus consisted of a yellow circle on a black screen presented in the center. The conditions were: no visual stimulus (No-V, listener-specific baseline), temporarily-synchronized audiovisual presentation (Sync) and asynchronous audiovisual presentation (Async). Sync and Async conditions corresponded to a measure of the audio-visual integration and distraction, respectively. All conditions were tested three times in a normalized Latin-square design. In order to eliminate hearing-loss as confounder, all subjects were normal hearing as assessed by pure-tone audiometry.

\textbf{Results}

For younger subjects (N=16), the MAAs in Sync were significantly (P<0.02) larger than in No-V. Preliminary results for elderly subjects (N=3) were not significantly different to that for younger subjects.

\textbf{Conclusion}

Stable MAAs, showing a significant ventriloquism effect, confirm the use of our procedure to investigate audiovisual interactions via headphones. Both elderly and young subjects showed similar baseline MAAs, demonstrating the suitability of our procedure for comparing different age groups. Similar MAAs were found in the integration and distraction conditions. The final results will be discussed in the light of the contribution of visual information in younger and elderly listeners.

The elderly can benefit from top-down repair: Aging and phonemic restoration

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\textbf{Background}

This study investigated the effect of age on the top-down repair of degraded speech using phonemic restoration of interrupted sentences.

\textbf{Methods}

Perception of meaningful sentences interrupted with silent intervals was measured with and without a filler noise in the gaps, with young (<26 years, average 22 years) and elderly (>62 years, average 66 years) normal-hearing listeners. The phonemic restoration effect, observed by an increase in intelligibility after the addition of the filler noise, provided a measure of top-down repair of interrupted speech. Further, the speaking rate of sentences was altered (by slowing down and speeding up by a factor of two) to observe the effects of cognitive processing speed on restoration.

\textbf{Results}

Consistent with literature, older adults performed worse than younger adults in general. However, despite the lower intelligibility scores with interrupted speech with silent gaps, once the filler noise was added, intelligibility reliably increased by large amounts. Hence, the elderly showed a remarkably robust restoration effect. Slowing down speech helped the
elderly adults gain more benefit than young adults with respect to phonemic restoration. This can be explained in terms of age-related cognitive slowing, as slowed speech gives elderly more time to extract cues from the speech and process them. Speeding up speech had a detrimental effect on speech intelligibility regardless of noise for both participant groups. This negative effect was larger for elderly than young adults, which can again be explained by cognitive slowing accompanying aging. With speeded speech there was a negative phonemic restoration effect, where inserting noise reduced the intelligibility of interrupted speech.

Conclusion
Even though elderly adults perform worse than young adults in understanding interrupted speech, they are still able to use top-down processes to extract cues and restore meaningful sentences when noise is added to the interruptions. It is conceivable that elderly rely on their life-long experience with language, as well as their rich vocabulary accumulated over a long time, to use these resources effectively to restore interrupted speech. When speech is slowed down such that the elderly are given more time to process the sentences, extract its cues and derive its content the restoration effect is strongly visible. When speech is speeded up, there is a detrimental effect of noise, producing very poor performance, as well as an reversed restoration effect. These results indicate that top-down repair could also be affected by age-related cognitive slowing.

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Endogenous Differences in Mitochondrial Metabolism Bias Sensory Cell Responses to Ototoxic Challenge
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Background
Aminoglycosides (AG), including gentamicin (GM), are the most frequently used antibiotics in the world despite the risk of irreversible cochlear damage and hearing loss (HL) associated with their use. High-frequency outer hair cells (OHCs) reliably generate and succumb to a barrage of AG-triggered pro-apoptotic signals, while inner hair cells (IHCs) display truncated pro-apoptotic signaling and greater survival rates. Although there are numerous causes of HL and deafness, reactive oxygen species (ROS) are now well-known instigators of multiple HL pathologies including: AG-induced ototoxicity, noise-induced and age-related HL. Given that ROS are normal byproducts of ATP synthesis that can rise to lethal levels when mitochondrial metabolism is perturbed, intrinsic differences in I/OHC mitochondrial metabolism may explain why high-frequency OHCs are profoundly sensitive to mitochondrial-mediated damage during various cochlear pathologies.

Methods
Mitochondrial metabolism was compared in apical, low-frequency (20% of cochlear length) and basal, high-frequency (80% of cochlear length) IHCs and OHCs obtained from acutely-cultured, intact cochlear (organ of Corti) explants harvested from mice (postnatal day 5 ±1d). Real-time changes in the reduced (NADH, fluorescent) and oxidized (NAD+, non-fluorescent) states of the metabolic intermediate, nicotinamide adenine dinucleotide, were used to measure both endogenous and GM-induced differences in IHC and OHC mitochondrial metabolism. NADH reduction and oxidation capacities were compared using sodium cyanide and FCCP, respectively.

Results
Despite similar quantities of functional mitochondria, endogenous NADH fluorescence was greatest in high-frequency OHCs. The dynamic range of NADH metabolism was also greatest in high-frequency OHCs (Left Fig.). Akin to adult cochlear preparations, GM rapidly decreased NADH fluorescence in high-frequency OHCs (Center Fig.). This decrease was shown to be due to a substantial decline in NADH reduction capacity rather than an increase in NADH oxidation (Right Fig.).

Conclusion
Basal turn, high-frequency OHCs are metabolically responsive a number of changes in their microenvironment including GM and elevated glucose concentrations. High-frequency IHCs and low-frequency IHCS and OHCs are substantially less sensitive to their microenvironment. The observed GM-induced decrease in mitochondrial metabolism in basal turn, high-frequency OHCs is consistent with the well-known enhanced susceptibility of high-frequency OHCs to undergo apoptosis after AG exposures. This metabolic predisposition may bias basal turn OHC responses to a variety of cochlear insults, particularly those involving energy metabolism and ROS production.
RNA-Sequencing Reveals the Involvement of the Complement Pathway in Noise-Induced Cochlear Damage

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Background

Exposure to intense noise causes sensory cell damage in the cochlea. Previous investigations have revealed multiple signal transduction pathways involved in cochlear degeneration, suggesting that noise-induced damage is a multifactorial process. To further explore novel signaling pathways, we examined the global transcriptome of noise-traumatized cochlear sensory epithelia using the next generation RNA-seq sequencing technique.

Methods

Young rats were exposed to a broadband noise at 120dB (SPL) for 2 h, and were sacrificed at 1 day post-noise exposure. The cochlear sensory epithelia were collected for total RNA isolation. The control animals received the same treatment except for the noise exposure. The samples from the two groups were used for cDNA library preparations and subsequently used for sequencing on the Illumina HiSeq 2000. Verification of the RNA-sequencing results was achieved using qRT-PCR and immunohistochemistry.

Results

Our sequencing results revealed an average of 176 ± 21 million reads for the normal control samples. Using a cut-off value of RPKM ≥ 0.1, we identified 12041 gene transcripts in the normal cochlear sensory epithelium. Comparing the RNA-seq data with the qRT-PCR data of selected genes examined in our previous investigation, we revealed a high correlation between the two data sets. In the cochlear samples collected 1 day after exposure to the broadband noise at 120dB (SPL) for 2 h, we found 164 ± 47 million reads, similar to the value seen in the normal cochleae. Using the cut-off value of RPKM ≥ 0.1, we identified 11874 gene transcripts. Among these gene transcripts, 71 were upregulated and 7 were downregulated. Using bioinformatic DAVID analysis, we found that these expression-altered genes are linked to multiple signaling pathways. Among these pathways, the complement pathway, an immune response-related pathway that participates in clearing dying cells from tissues, has not been implicated in cochlear pathogenesis. Further qRT-PCR and immunolabeling assessments confirmed that the two genes (C1s and Cfi) identified by RNA-seq for the complement pathway were indeed involved in the pathogenesis of the organ of Corti. Importantly, the change in Cfi was associated with sensory cell death.

Conclusion

Identification of C1s and Cfi in the cochlear sensory epithelium following acoustic trauma highlights the presence of a cochlear inflammatory response, which may be modulated towards reducing the extent of sensory cell damage. (Supported by NIH R01 DC010154-01A2 to BH Hu).

A Novel Treatment Reduces Hearing Loss, and Cochlear and Brain Injury after Noise and Blast

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Background

High level steady state noise and blast can lead to cochlear injury and hearing loss and stress and/or injury to brainstem and cortical and other auditory centers. Military personnel exposed to blast can experience, hearing loss, tinnitus, and other associated co-morbidities related to memory and orientation impairment for example. Because noise-induced
hearing loss and tinnitus continue to be ongoing prevalent problems despite traditional hearing conservation programs, research is being pursued to achieve a pharmacological treatment for these injuries. Here we report that a novel pharmacological therapy reduces cochlear and brain stress/injury and permanent hearing loss induced by high level steady state noise as well as blast.

**Methods**

Male Long-Evans pigmented rats had baseline ABR and DPOAE testing followed by exposure to either 115 dB SPL narrow band noise for 1 hour or 3 or 4, 14 p.s.i. blasts. One hour after the noise/blast exposure rats were treated with an intraperitoneal injection of a nitrone (HPN-07) plus the glutathione prodrug n-acetylcysteine (NAC), or, with solution vehicle only. Serial ABR and DPOAE measurements were made up to 21 days post exposure. At different time points animals were euthanized and cochlear and brain tissues were harvested and processed for histological and immunohistochemical analysis.

**Results**

Treatment substantially reduced permanent hearing loss induced by narrow band noise or blast. Inner and outer hair cell losses were also substantially reduced after noise or blast trauma with treatment. Treatment also reduced injury biomarker expression in the brainstem, hippocampus, and auditory cortex after blast exposure. After noise exposure, treatment reduced c-fos expression in the cochlear nucleus.

**Conclusion**

These data suggest it may be possible to develop a pharmacological approach that not only reduces blast-and noise-induced cochlear damage and hearing loss but reduces brain injury associated with a blast and the stress response in the cochlear nucleus associated with high level noise exposure. This could have implications for noise or blast induced tinnitus. It may also have implications for the treatment of mild traumatic brain injury associated with blast in military personnel. Further research is needed to establish the extent of brain injury recovery and tinnitus reduction that might be possible with this treatment approach. A single pharmacological treatment that addressed both cochlear injury as well as brain injury associated with high level noise or blast could be useful in treating several co-morbid conditions experienced by military personnel.
Subcutaneous Etanercept Attenuates Noise-Induced Hearing Loss
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Background
Noise induced hearing loss (NIHL) is the most common cause of hearing loss in adults. Our laboratories as well as others have shown that noise exposure induces reactive oxygen species (ROS) generation in the cochlea which initiates an inflammatory response. We have shown that noise exposure increases the expression of pro-inflammatory mediators such as tumor necrosis factor-α (TNF-α), inducible nitric oxide synthase (iNOS) and cyclooxygenase-2 (COX2), and increased CD14 positive cells in the rat cochlea. Therefore, we hypothesized that inhibition of the inflammation could be a potential therapeutic strategy for treating NIHL. In this study, we tested the effectiveness of the TNF-α receptor inhibitor (Etanercept - ETA) against NIHL.

Methods
To determine the utility of ETA to protect hearing, baseline ABR testing was recorded. Rats were then administered vehicle (saline) or ETA by the transtympanic (TT) (250µg in 50µl saline) or subcutaneous (SC) (2.5 mg/kg) routes, followed by noise exposures at 3 and 7 days following ETA treatments. To determine the efficacy of ETA against noise trauma, rats were treated with saline (via the IT or SC routes) or ETA at 24 or 48h following noise exposure. Noise was administered at 122 dB OBN (centered at 16 kHz) for 1 h. Post-treatment ABRs were recorded 21 days after noise exposure. Cochleae were harvested for morphological analysis by scanning electron microscopy (SEM), immunohistochemistry and gene expression studies.

Results
Rats treated with ETA (SC route) 2h following noise showed almost complete protection from NIHL (<5dB ABR shift) and significant protection up to 24h. ETA administration (TT route) provided significant protection at 16 and 32 kHz (<25dB ABR shift). Administration of ETA 7d prior to noise exposures provided significant protection from hearing loss. This Otoprotection afforded by ETA was associated with preservation of outer hair cells, decreased gene expression of stress-response genes, such as NOX3 NADPH oxidase and TRPV1 and pro-inflammatory mediators such as TNF-α, iNOS and COX2.

Conclusion
ETA significantly reduced noise induced hearing loss when administered within 2h-24h after noise exposure. It was also shown to mediate otoprotection when administered up to 7days prior to noise exposure. These findings support the clinical use of etanercept (FDA approved) to prevent and treat NIHL. (Supported by HHF).

Expression of Inflammatory Cytokines and Ion Homeostasis Genes in the Cochlea Is Up-regulated by Lipopolysaccharide-induced Sepsis
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Background
The mechanism by which sepsis-induced inflammation enhances aminoglycoside trafficking into the cochlea remains poorly understood. This study sought to determine if sepsis and/or aminoglycosides induced changes in gene and protein expression of inflammatory-mediated cytokines and ion homeostasis proteins in the cochlea.

Methods
Three groups of mice received one of the following treatments: systemic lipopolysaccharides (LPS) for 6 hours for sepsis induction, systemic gentamicin for 3 hours, or both LPS and then gentamicin treatment. A fourth group served as controls. After sacrifice, cochleae were collected and underwent either RNA extraction for PCR, or protein extraction for enzyme-linked immunosorbent assay (ELISA). A total of 32 cytokine and ion homeostasis genes/proteins were analyzed.

Results
Cochlear gene expression analysis revealed that gentamicin alone did not induce significant inflammation or other changes in the cochlea within 3 hours. However, significant inflammatory and ion homeostasis gene up-regulation was seen with bacteria-derived LPS treatment. The most up-regulated cytokine genes with LPS were interleukin (IL)-6, macrophage inflammatory protein (MIP)-2α, and neutrophil activating protein (CXCL)-1. In the cochlear protein quantification experiments, the cytokines found in greatest quantity were IL-6, MIP-2α, and CXCL1. A similar distribution was observed when measuring the cytokines in the serum. The ion homeostasis genes that LPS up-regulated in a
biologically significant manner (fold change ≥ 1.5) were gap junction-α1, claudin-4, and claudin-14. Gentamicin treatment did not modulate these LPS-induce changes.

**Conclusion**

Sepsis induces a systemic pro-inflammatory state that can enhance aminoglycoside uptake by cochlear tissues. After LPS administration in mice, which induces systemic inflammation, there is statistically significant up-regulation of cytokine gene and protein expression, and of specific ion homeostasis genes, in the cochlea regardless of aminoglycoside exposure. Interestingly, the most significant cytokines found in the gene expression analysis were also those seen in the protein quantification experiments. Given that cytokines increase permeability across blood-endothelial barriers, this phenomenon could enhance aminoglycoside trafficking into cochlear tissues. LPS-induced up-regulation of ion homeostasis gene expression may provide additional clues to the mechanisms by which enhanced aminoglycoside trafficking into the cochlea during systemic infection occurs. Thus, systemic infection may synergistically enhance the cochlear uptake of aminoglycosides and subsequent ototoxicity.

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**Inner ear changes in Niemann-Pick disease type C (NPC), and in NPC therapy, in the mouse.**

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**Background**

Niemann-Pick disease type C (NPC) is a rare autosomal recessive lipid storage disorder that affects lysosomal transport of cholesterol and results in progressive neural and visceral organ dysfunction and eventual death. The central auditory system has been shown to be affected in NPC. Peripheral auditory system effects are reflected in reduced distortion product otoacoustic emissions (DPOAEs). We studied morphological changes in the NPC mouse inner ear. The FDA approved 2-hydroxpropyl-beta-cyclodextrin (Cyclo) for treatment in NPC patients in 2009, under compassionate use status. At that time, Cyclo had been shown to effectively alleviate symptoms and increase life expectancy in mice. Initial results in humans are promising. However, Cyclo treatment in cats causes rapid and irreversible hearing loss. A better understanding of the development of hearing loss in the peripheral auditory system in NPC, with or without the administration of Cyclo, could lead to treatment advancements. Therefore we also studied the ototoxicity of Cyclo in mice.

**Methods**

Adult mice were given either a high single or low repeated doses of Cyclo(1,000-8,000 mg/kg body weight, subcutaneously). ABRs were recorded at 4, 16, and 32 kHz before and after treatment. Inner ear tissue from normal, Cyclo- treated and BALB/c-npc1NIH mice was processed and sectioned for light and electron microscopic viewing. The opposite ears were micro-dissected for organ of Corti analysis.

**Results**

Changes were seen in NPC mice at the stria vascularis, nerve endings in the organ of Corti and in the spiral ganglion. Cyclo administration caused significant ABR threshold shifts at 4 kHz and 16 kHz 3 weeks following a high single dose. Less robust changes were seen in the mice given lower repeat doses. These results were also reflected in the organ of Corti pathology.

**Conclusion**

Neural changes seen in the present study are consistent with damage in the central nervous system that is known to occur in NPC. Surprisingly, while the hair cells require cholesterol for normal functioning, they appear to be relatively unaffected. The DPOAE changes are more likely due to strial damage. The changes seen after Cyclo treatment appear to be related to the loss of membrane cholesterol in the outer hair cells. New treatments are being tested and will be discussed. (Funding: Ara Parseghian Medical Research Foundation (JJR) & Pharmacology Training Grant at UT Southwestern, NIH T32 GM007062 (Mangelsdorf))
Molecular Mechanisms of the PRPS1 Deafness
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Background
PRPP (phosphoribosylpyrophosphate) synthetases (PRPPs) are a family of enzymes that catalyse the synthesis of PRPP, a substrate in and an allosteric regulator of nucleotide synthesis. *PRPS1* loss-of-function missense mutations cause X-linked nonsyndromic sensorineural deafness type 2 (DFNX1) (Liu et al. 2010) or syndromic deafness including Arts syndrome and X-linked Charcot-Marie-Tooth disease-5 (CMTX5). However, the molecular mechanisms of the enzymatic aberration in DFNX1, as well as the biological effects of *PRPS1* mutations, are not understood.

Methods
To investigate the pathogenic mechanism of the PRPS1 deafness associated mutations, we have used the antisense morpholino (MO) technology to knock down *Prps1* gene function in zebrafish.

Results
By RT-PCR, we found that injection with M3, an acceptor site-targeting MO in *Prps1b* into zebrafish embryos, results in production of aberrantly spliced message mRNA. Our development study shows that there is a delayed hatching in M3 morphants and they have abnormalities in both anterior and posterior otoliths. Measurements of larval zebrafish inner ear and of saccular otolith growth indicates that the total inner ear and saccular otolith area of larval zebrafish injected with M3 are smaller compared with those measured in controls. Furthermore measurements of microphonic potentials revealed that the microphonic responses from the morphant's ears were considerably reduced compared to those measured in controls, indicating that these morphants had impaired mechanoelectrical transduction.

Conclusion
Overall, our findings emphasize the need for tight regulation of PRPS1 activity and imbalances in nucleotide metabolism affect nearly the whole cellular function. As the metabolic enzyme critical for nucleotide biosynthesis, PRPS1 is a promising target for drug in prevention and treatment of hearing loss.

Noise exacerbates Hypoxia-associated changes in the Alport Mouse Cochlea
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Background
Mice modeling the recessive form of Alport Syndrome have thickened strial capillary basement membranes (SCBMs). The cochlea of Alport mice is inherently hypoxic with significant differences post-noise exposure in ABR thresholds, EP magnitude and Na,K-ATPase activity compared to wild-type (WT) littermates. Our previous studies show Alport mice exhibit increased heparan sulfate proteoglycan in SCBMs and upregulation of hypoxia-regulated genes/proteins (i.e. facilitated glucose transporter-1, vascular endothelial growth factor) in the cochlea. These data suggest that reduced transcapillary movement of oxygen, metabolic nutrients and waste due to altered SCBM filtration properties creates a hypoxic microenvironment in scala media. We hypothesize that the combination of thickened SCBMs and metabolic stress drives the Alport cochlea into a glycolytic state to meet ATP demand for cellular metabolism and auditory transduction.

This project aimed to quantify energy states and oxidative stress in the cochlear lateral wall (LW). Levels of LW intracellular ATP and total glutathione, including oxidized to reduced GSSG/GSH ratio, were measured. To further characterize the biochemical dysregulation occurring in this hypoxic microenvironment, these assays were conducted after exposure to metabolic stress.

Methods
Lateral wall tissue was microdissected from 7-9 week Alport mice and their WT littermates. A subset of each genotype was subjected to noise (10 kHz OBN, 120 dB SPL, 10h). The LW was homogenized and pelleted by centrifugation. The supernatant was assayed for ATP and glutathione levels by flash/ glow luminometry. Data was referenced to protein concentration.

Results
Inherent metabolic differences were observed post-noise between WT and Alport mice. From quiet to post-noise, the WT LW has a robust, aerobic increase in intracellular ATP, denovo total glutathione (GSH+GSH) synthesis and maintenance
of GSSG/GSH ratio. In contrast, the post-noise Alport LW intracellular ATP level remains unchanged, with only a modest increase in total glutathione and a substantial increase in GSSG/GSH ratio.

Conclusion
The current study confirms and extends our previous work demonstrating that the Alport LW is essentially hypoxic, with an inability to efficiently regulate intracellular ATP and maintain GSSG/GSH ratio post-noise exposure. This implies that oxidative stress/free radical formation from mitochondrial dysfunction underlies the decreased physiological measures (ABR, EP, Na,K-ATPase) in the Alport cochlea. These results prompt further investigation of mitochondrial respiration through studying energy charge balance (e.g. ADP/ATP ratio), ROS formation, as well as glycolytic respiration (e.g. pyruvate/ lactate ratio).

Reactive oxygen species generation and detection in the stria vascularis of Alport mice

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Background
Alport syndrome is a hereditary defect in type IV collagen genes. Mice modeling the recessive form of Alport syndrome demonstrate progressive thickening of strial capillary basement membranes (SCBM), a mild high frequency hearing loss and sensitivity to noise. Our previous studies show that the cochlea is hypoxic and exhibits upregulation of hypoxia-regulated genes/proteins (i.e. HIF-1α, GLUT-1, VEGF). Noise-exposed Alport mice are unable to modify intracellular ATP levels, resulting in increased oxidized to reduced glutathione (GSSG/GSH) ratio, indicating disrupted mitochondrial respiration. Mitochondrial dysfunction, with loss of redox homeostasis and formation of excessive free radicals, may play an important role in the pathogenesis of the hearing loss associated with Alport syndrome. A method for detecting/quantifying ROS formation in the stria could be utilized to examine whether thickened strial capillary basement membranes result in mitochondrial dysfunction under conditions of metabolic stress.

This project sought to:
-- develop a protocol using the fluorescent marker MitoSox (Invitrogen), an indicator of mitochondrial ROS generation in vitro, to measure oxidative stress in vivo.
-- determine the level of reactive oxygen species generation in Alport stria vascularis with and without metabolic stress.

Methods
Cochlea lateral wall (LW) tissue was microdissected from 7-9 week old Alport mice and their WT littermates. A subset of each genotype was subjected to noise (10 kHz OBN, 120 dB SPL, 10h). The experimental LW was incubated (0.625M MitoSox, 10min) prior to mounting flat, stria upward, using DAPI gold. The contralateral LW served as a control, mounted directly in DAPI Gold. Slides were kept at 4°C until visualization. The total time for dissection, incubation and mounting was ≤ 35 minutes. The LW was visualized with confocal microscopy. Two-channel z-stacks, ROS and nuclear stain, were analyzed with Volocity software to quantify non-nuclear ROS.

Results
-- Protocol developed to quantify ROS using confocal microscopy to detect MitoSox in unfixed, isolated cochlear tissue.
-- Staining patterns of MitoSox correlate to expected areas of ROS formation in the stria vascularis based on predicted areas of greatest mitochondrial density.
-- WT and Alport mice in quiet demonstrate similar ROS levels.
-- Following noise exposure, ROS levels differ between the WT and Alport mice.

Conclusion
-- ROS formation in the stria vascularis can be reliably imaged and quantified.
-- Mice exhibiting thickened SCBMs and cochlear hypoxia demonstrate mitochondrial dysfunction under conditions of metabolic stress.
-- Mitochondrial dysfunction contributes to the pathogenesis of the hearing loss associated with Alport syndrome.
IGF-1 deficit predisposes to noise-induced hearing loss in mice.
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Background
Human IGF-1 deficiency (ORPHA73272, OMIM608747) is a rare disease associated with poor growth rates, mental retardation and syndromic hearing loss. Equally, Igf1⁻/⁻ mice are dwarfs with poor survival rates and congenital profound deafness¹. IGF-1 is a neuroprotective agent, and accordingly, the physiological age-related decrease in circulating IGF-1 levels have been related to cognitive and brain alterations. Igf1⁻/⁻ mice present undetectable serum levels of IGF-1 throughout its life, whereas Igf1⁺/⁻ and Igf1⁺/+ littermates show an age-dependent decrease in IGF-1 serum levels, especially from 6 months of age on², ³, which correlates with the increase in ABR thresholds. However, there is little information on the effect of low levels of IGF-1 on the susceptibility to noise induced hearing loss (NIHL) and the potential protective actions of IGF-1 against this condition.

Methods
We have studied the susceptibility of Igf1⁺/⁻ and Igf1⁺/+ mice to NIHL at different ages, with functional (auditory brainstem responses, ABR), morphological (cochlear histology and stereological hair-cell quantification) and molecular (RT-qPCR, Western Blotting) studies.

Results
Noise-exposure experiments with 1 and 3 months-old mice did not reveal differences between genotypes, both genotypes were equally sensible to NIHL. However, 6 month-old Igf1⁺/⁻ presented a greater susceptibility to noise damage, with higher threshold shifts and a poorer recovery compared to noise-exposed Igf1⁺/+ mice. Igf1⁺/⁻ mice showed severe morphological changes, including the massive loss of hair cells, as well as alterations in the main IGF-1 signalling pathways. These changes correlated with low IGF-1 serum levels in the heterozygous mice, compared to wild type.

Conclusion
These results support the idea that low levels of IGF-1 predisposes to higher susceptibility to NIHL. Therefore, IGF-1-based therapies could contribute to prevent or ameliorate age related and noise-induced hearing loss.

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See also communication num. 33
Preliminary Results of Treating Deafness in Connexin Knockout Mice Using Retinoids
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Background
Connexin mutations represent the most common cause of nonsyndromic hereditary deafness, with 10-50% of all patients in different ethnic populations bearing mutations specifically in the GJB2 gene, which encodes the connexin (Cx) 26. Previous studies have shown the promise of treating nonsyndromic hereditary deafness caused by Cx mutations by increasing expressions of the affected Cx or even the remaining Cx levels (e.g. Cx26 in the absence of Cx30). Retinoids, the family of vitamin A derivatives, have been utilized to increase expression of the Cx levels (e.g. Cxs26, 30 and 43) in dermatological and oncological studies, some of which involve syndromic deafness conditions caused by the Cx mutations such as keratitis-ichthyosis-deafness (KID) syndrome and palmoplantar keratoderma with deafness. Moreover, retinoids have been used to treat various types of hearing loss in both clinical and experimental applications, with promising results in vivo and in vitro. Yet the impact of retinoids on Cx expression in the inner ear is largely unknown.

Methods
In this study we examined the impact of retinoid derivatives in the mouse inner ear by surgically implanting gel foams soaked in the retinoids (e.g. retinyl palmitate and retinyl acetate) at the round window on postnatal day 1 (P1). At P10, the cochlea was harvested and examined for morphological changes and the expression of Cx26 and Cx30.

Results
Application of retinyl acetate greatly reduced the Cx30 expression, at concentrations 30 μM and greater, to the level below that detectable by immunobolabeling. For the expression of Cx26, there was a concentration dependent decrease assayed by immunolabeling. Furthermore, application of retinyl palmitate dramatically affected the development of the sensory epithelium, causing the tunnel of Corti to stay in a closed state. The mechanism underlying such a profound morphological change may also point to the modulation of the gene expression of cochlear Cxs, as seen in the conditional Cx26 null mouse models we previously reported. Further studies will be conducted to investigate the gene expressions in response to applications of retinoids, the impact on hearing, and to determine the mechanism underlying the closure of the tunnel of Corti.

Conclusion
Our findings point to a delicate relationship between different Cxs and retinoids in the inner ear development, and may be an important step in fully understanding the underlying mechanisms of retinoids in treating hearing loss.
Connexin26 Mutations that Cause Hereditary Deafness Lead to Macromolecular Complex Degradation of Cochlear Gap Junction Plaques
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Background
Hereditary deafness affects about 1 in 2000 children and GJB2 gene mutation is most frequent cause for this disease in the world. GJB2 encodes connexin26 (Cx26), a component in cochlear gap junction.

Methods
In this study, we analyzed macromolecular change of gap junction plaques with two different types of Cx26 mutation as major classification of clinical case, one is a model of dominant negative type, Cx26R75W+ and the other is conditional gene deficient mouse, Cx26f/fP0Cre as a model for insufficiency of gap junction protein.

Results
Gap junction composed mainly of Cx26 and Cx30 in wild type mice formed large planar gap junction plaques (GJP) along the cell junction site with the adjacent cells in normal inner sulcus cells and border cells. In contrast, Cx26R75W+ and Cx26f/fP0Cre showed fragmented small round GJPs around the cell junction site and marked decrease of total gap junction area. In Cx26f/fP0Cre, some of the cells with Cx26 expression due to their cellular mosaicism showed normal large GJP with Cx26 and Cx30 only at the cell junction site between two Cx26 positive cells. These indicate that bilateral Cx26 expressions from both adjacent cells are essential for the formation of the cochlear linear GJP, and it is not compensated by other cochlear Connexins such as Connexin30.

Conclusion
In this study, we demonstrated a new molecular pathology in most common hereditary deafness with different types of Connexin26 mutations, and this machinery can be a new target for drug design of hereditary deafness.
Morphological Changes of Cochlear Gap Junction Plaques in Brn4 Deficient Mouse, a Mouse Model of DFN3 Non-syndromic Deafness

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Background
Brn4, which encodes a POU transcription factor, is the gene responsible for DFN3, an X chromosomelinked nonsyndromic hearing loss. Brn4 deficient (KO) mice show low endocochlear potential (EP), hearing loss and severe ultrastructural alterations in spiral ligament fibrocytes. Minowa O et al., Science, 1999. In humans, mutations in the connexin26 (Cx26) and connexin 30 (Cx30) genes, which encode gap junction proteins and are expressed in cochlear fibrocytes and nonsensory epithelial cells (cochlear supporting cells) to maintain proper EP, are well known to be responsible for hereditary sensorineural deafness. The molecular pathologies of Brn4 deficiency causing low EP are still unclear.

Methods
It has been hypothesized that gap junction in the cochlea provide an intercellular passage by which K+ are transported to maintain high levels of the endocochlear potential essential for sensory hair cell excitation. In this study, we analyzed the formation of gap junction plaques in cochlear supporting cells of Brn4 KO mice in different stages by confocal microscopy and three dimensional graphic construction.

Results
Gap junction composed of mainly of Cx26 and Cx30 in wild type mice showed horizontal linear gap junction plaques along the cell-cell junction site with the adjacent cells and these formed pentagonal or hexagonal outlines of normal inner sulcus cells and border cells. However, the gap-junction plaques in Brn4 KO mice did not show normal linear structure, but the round small spots were observed around the cell-cell junction site. The size of gap junction units of Brn4 KO mice was significantly shorter than control mice.

Conclusion
Our results demonstrated that Brn4 gene mutation affects on the accumulation and localization of gap junction proteins at the cell border among cochlear supporting cells. It may suggest that Brn4 significantly associates with cochlear gap junction properties to maintain proper EP in cochlea as well as the mutations of Cx26 or Cx30.

Evaluation of Auditory Function in Slc44a2 Mutants on an ARHL-resistant Genetic Background

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Background
SLC44A2 (CTL2, choline transporter-like protein 2), is a target of antibody-induced hearing loss and carries the HNA3a, 3b antigen of transfusion related acute lung injury (TRALI). We developed a knockout mouse to investigate the function of SLIC44A2 protein in the cell border among cochlear supporting cells. It may suggest that Brn4 significantly associates with cochlear gap junction properties to maintain proper EP in cochlea as well as the mutations of Cx26 or Cx30.

Methods
We backcrossed B6.129-Slc44a2del-ex3-10/Tec to FVB mice, which carry the ARHL-resistant Cdh23753G allele at Ahl, then genotyped subsequent intercross progeny for the Cdh23753G variant and for the Slc44a2 deletion using PCR-based assays. Intercross progeny were tested at 1.5 months and 4.5 months of age for hearing in this genetic context.

Results
Eighteen intercross mice were genotyped and evaluated for auditory function. Three mice were Slc44a2del-ex3-10/del-ex3-10 Cdh23753G/G, two were Slc44a2del-ex3-10/del-ex3-10 Cdh23753G/A, two were Slc44a2+/+ Cdh23753G/G and two were Slc44a2+/ Cdh23753A/G. Eight mice were Slc44a2del-ex3-10/+ Cdh23753G/G and one was Slc44a2del-ex3-10/+ Cdh23753A/G. Progressive hearing loss at 48kHz occurred in three mice with homozygous Slc44a2 exon 3-10 deletion and wild type Cdh23753G/G (75, 73, 60dB) and two with homozygous Slc44a2 exon 3-10 deletion heterozygous for Cdh23753A/G had hearing loss(35, 105dB). Of four mice homozygous for wt Slc44a2, two with Cdh23753G/G and one with Cdh23753A/G had no
hearing loss, but one with \( \text{Cdh23}^{53G/A} \) had a high frequency hearing loss (48 kHz, 60dB SPL). Of nine mice heterozygous for \( \text{Slc44a2}^{\text{del-ex3-10/a}} \), six had high frequency hearing loss (45-80db SPL) five with \( \text{Cdh23}^{53G/G} \), and one with \( \text{Cdh23}^{53G/A} \) had normal hearing.

**Conclusion**

Significant high frequency hearing loss occurred by 4.5 months in nearly all mice carrying the \( \text{Slc44a2} \) exon 3-10 deletion, even in those carrying an ARHL-resistant allele of \( \text{Ahl} \). This result is consistent with a role for the \( \text{Slc44a2} \) gene in normal auditory function.

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**Hearing Threshold Differences among C57BL/6J (wild-type), FVB/NJ (wild-type), and \( \text{Cdh23}^{\text{ahl}} \) corrected FVB.B6.129 mice**

_Bala Naveen Kakaraparthi*, Irina Laczkovich*, Thankam Nair, Pavan Kommareddi, Danielle Miller, Lisa Kabara, Ariane Kanicki, David Dolan, David Kohrman, Thomas Carey*

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**Background**

C57BL/6J mice and embryonic stem cells from 129S/129X1 strain, used to develop knockout strains, share a \( \text{cadherin23} \) G->A mutation at codon 753 designated \( \text{Ahl1} \) or \( \text{Cdh23}^{\text{ahl}} \). This creates problems for hearing research due to early-onset presbycusis detectable at high frequencies (HF) from 1.5 months of age. We developed a \( \text{SLC44A2} \) (CTL2, choline transporter-like protein 2), which is a target of antibody-induced hearing loss, knockout mouse using B6.129 strain to examine CTL2 function. The knockout mouse exhibited HF hearing loss, but because the background strain was a confounder, we backcrossed to FVB/NJ (\( \text{Cdh23}^{+/+} \)). We compared hearing in wild type (wt) C57BL/6J, FVB/NJ, and FVB.B6.129.

**Methods**

We backcrossed B6.129 homozygous for \( \text{Cdh23}^{\text{ahl/ahl}} \) to FVB/NJ, wt for \( \text{cadherin23} \), and tested offspring for the \( \text{Cdh23}^{\text{ahl/ahl}} \) mutation. After a second backcross to FVB/NJ, the mice were interbred and tested for the \( \text{Ahl1} \) polymorphism (\( \text{CGGTGAG vs. CCGTGGAG} \)) by PCR and Mspl digestion, which yields a 600bp band from the mutant (\( \text{CCAG} \)) and two bands of 200 and 400bp from the wt allele (\( \text{CCGG} \)). We compared the wt (\( \text{Cdh23}^{+/+} \)) FVB.B6.129 mice without the \( \text{Ahl} \) mutation to wt C57BL/6J (\( \text{Cdh23}^{\text{ahl/ahl}} \)) mice and FVB/NJ (\( \text{Cdh23}^{+/+} \)) controls by testing their auditory brainstem response (ABR) at 12, 24, and 48 kHz at 1.5 months, 3 months, and 4.5 months of age.

**Results**

Sixteen C57BL/6J mice were tested, and some were euthanized at intervals to examine ears for inner ear morphology. At 1.5 months old, the average hearing threshold at 48 kHz was ~59.5dB (thresholds > 50 dB translate to hearing loss). Progressive HF hearing loss occurred at 3 months (~87.5dB) and 4.5 months of age (109dB). Three FVB/NJ controls had averages of 30dB at 1.5 months and ~36dB at 3 months at 48 kHz. The two \( \text{Cdh23}^{\text{ahl}} \) corrected mice (\( \text{Cdh23}^{+/+} \)) also exhibited normal hearing, with averages of 25dB at 1.5 months and 20dB at 4.5 months at 48 kHz.

**Conclusion**

C57BL/6J mice exhibit progressive HF hearing loss from 1.5 months onward. FVB/NJ backcrossing and \( \text{Cdh23}^{\text{ahl}} \) correction resulted in normal hearing thresholds that remained constant up to 4.5 months of age. The results demonstrate that correction of the \( \text{Ahl1} \) mutation in the C57BL/6J and 129 strains is necessary for knockout experiments of genes involved in hearing acuity.

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**Examining the Role of Gtf2ird1 Deletion in Auditory Physiology and Behavior in a Murine Model of Williams-Beuren Syndrome**

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**Background**

Williams-Beuren Syndrome (WBS) is a genetic disorder typified by a constellation of features, including craniofacial and neuro-developmental abnormalities, which have been investigated substantially. However, less is known about the etiologies of auditory abnormalities associated with WBS such as hyperacusis, tinnitus, and high-frequency hearing loss.
Here we examine auditory physiological and behavioral characteristics of mice with deletion of Gtf2ird1, one of the 26 genes located within the WBS region of chromosome 7, that display neuro-developmental features characteristic of WBS.

Methods
Mice that were homozygous knock-out (KO) for the Gtf2ird1 gene were compared with heterozygous (HET) and wild-type (WT) littermates on measures of auditory function. Auditory brainstem response (ABR) and distortion product otoacoustic emissions (DPOAEs) were measured to assess cochlear and retro-cochlear function in young (~8wks) mice, as well as in older (~17wks) mice that were noise-exposed (8-16 kHz, 100 dB SPL, 2 hrs) at 6wks. The same groups of mice underwent acoustic startle response (ASR) and prepulse inhibition (PPI) of startle tests to assess supra-threshold auditory behavior.

Results
In the young cohort, KO and HET mice showed slight elevation of ABR and DPOAE thresholds compared to WT littermates, especially for high (> 22 kHz) frequencies. ABR waveform morphology was comparable across genotypes with the exception of a larger amplitude wave 2 in KO and HET mice compared to WT for supra-threshold, mid-frequency (8-22 kHz) stimuli. Behavioral responses to 16 kHz prepulses and to broadband startle stimuli were similar across genotypes. In the older, noise-exposed cohort, ABRs of KO and HET mice diverged, with KO mice showing lower thresholds and greater preservation of ABR wave amplitudes.

Conclusion
This study is one of the first to examine auditory function and behavior in a mouse model of WBS. Young mice with homozygous or heterozygous deletion of Gtf2ird1 show subtle auditory threshold elevation, similar to some individuals with WBS. Reflex-based tests of auditory function reveal no significant behavioral differences and no obvious indication of hyperacusis-like behavior. Interestingly, in an older cohort of mice with history of acoustic overexposure, KO mice showed greater preservation of thresholds and supra-threshold neural response to sound compared to HET littermates. Taken together with the known expression of Gtf2ird1 in the adult murine cochlea, further investigation is warranted to understand the role of this gene in the normal and injured ear.
A Mouse Model of Progressive Bone Remodeling and Mixed Hearing Loss
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Background
Otosclerosis is a disease of the otic capsule that is among the most common causes of acquired hearing loss. We have previously demonstrated that the osteoprotegerin (OPG) knockout mouse exhibits active bone remodeling within the otic capsule and progressive, apparently mixed (conductive and sensorineural) hearing loss (Zehnder et al 2006), characteristics seen in otosclerosis. Both the bone remodeling and the threshold elevations are responsive to treatment with bisphosphonates (Adachi et al 2008). OPG mutants used in these studies, however, existed in C57BL/6J (B6), a strain with rapidly progressive, age-related hearing loss (AHL). We have now bred the mutation into the normal hearing, normally aging CBA/CaJ strain, allowing for better assessment of the nature of the hearing loss attributable to OPG deficiency and the long-term efficacy of drug treatment. Here, we describe the age-progressive auditory pathophysiology and histopathology of the OPG mutant in the new background.

Methods
Experiments were conducted on an age-graded series (4 to 64 wk) of OPG-deficient (-/-, +/-) mice and wildtype (+/+ ) littermates in the 10th generation of transfer to CBA/CaJ. Animals underwent auditory function testing using DPOAEs and ABRs (5.6 – 44.06 kHz). Paraffin-embedded middle ear and cochlear tissues (H&E stained) were examined by light microscopy to characterize the histopathology.

Results
OPG -/- (but no +/- or +/+) mice demonstrate abnormal remodeling of bone within the otic capsule. Foci show osteoclastic bone resorption and formation of new bone, and are progressive in number, size and location as animals age. Marrow spaces in bone are dramatically enlarged. Functional assays reveal progressive hearing loss in -/- compared to age-matched +/- and +/- mice. Mild/moderate high frequency loss by ABR at 4 weeks progresses to severe loss across frequency by 32 weeks. DPOAE losses exceed those by ABR before limitation by a lower response ceiling. This asymmetry suggests conductive involvement affecting both forward and backward transmission through the middle ear, producing compound loss in the DPOAEs. Heterozygotes are wildtype-like in structure and function at all ages.

Conclusion
The OPG -/- mouse in CBA/CaJ shows progressive otic capsule bone remodeling and hearing loss without the complication of B6-related AHL. Studies aimed at clarifying the relative hearing loss consequences of the middle ear vs. cochlear sites of lesion are underway, to support our efforts to characterize long-term effects of drugs with potential efficacy in the treatment of otosclerosis. Research supported by NIDCD: RO1 DC03401

Restoration of Hearing in VGLUT3 Knock Out Mice using Virally-Mediated Gene Therapy (P1-3 Injection)
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Background
Mice lacking the vesicular glutamate transporter 3 (VGLUT3 KO) are born deaf due to the absence of glutamate release from the inner hair cells. In a prior study, we demonstrated correction of this genetic defect through virally-mediated gene therapy, using an in vivo microinjection of adeno-associated virus with a VGLUT3 insert (AAV-VGLUT3) into the endolymph of P10-12 VGLUT3 KO mice.

Methods
In the present study, in vivo microinjection of AAV-VGLUT3 via direct injection through the round window membrane was performed on P1-3 VGLUT3 KO mice.

Results
ABR testing demonstrated restoration of hearing in all VGLUT3 KO mice, to thresholds equivalent to those seen in wild-type mice. Long-term follow up showed that in the P1-3 rescued mice, 100% of animals had preservation of normal ABR thresholds through 9 months. In contrast, in mice rescued at P10-12, only 11% of animals demonstrated hearing
preservation at this same time point. Immunofluorescence with anti-VGLUT3 antibody demonstrated robust VGLUT3 expression (100% of IHC transfected) within the inner hairs cells of the rescued VGLUT3 KO mice. In contrast, in mice rescued at P10-12, only 40% of IHC were transfected using the same amount of virus. Lastly we noticed an improved startle-response to hand clap of the P1-3 injected as compared to the P10-12 rescued mice. However, reduced spiral ganglion cell counts that are seen in the KO mouse were not reversed in the P1-3 rescued mice.

Conclusion
The results of this study again demonstrate functional recovery of hearing in a genetic model of human deafness due to loss of VGLUT3 function, using virally mediated gene therapy. Furthermore, earlier injection of the virus showed higher efficacy in hearing recovery, normalization of hearing to wild type ABR levels, greater longevity of the hearing response and better startle to hand clap than the mice rescued at P10-12. However, there was still no difference in the spiral ganglion cell counts between KO and rescued mice, suggesting that pre-natal spontaneous inner hair cell release of glutamate helps maintain normal spiral ganglion cell numbers through birth.

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Otopathology in Congenital Toxoplasmosis
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Background
Toxoplasmosis is a parasitic infection caused by Toxoplasma gondii. If fetal infection occurs early in gestation severe inflammation and necrosis can cause brain lesions, chorioretinitis and hearing loss. Hearing loss in congenital toxoplasmosis may be preventable with early diagnosis and treatment.

Methods
The temporal bones of nine subjects with congenital toxoplasmosis were removed at autopsy and studied under light microscopy. Cytocochleograms were constructed for hair cells, the stria vascularis and cochlear neuronal cells.

Results
Three of nine subjects (33%) were found to have parasites in the temporal bone. The organism was identified in the internal auditory canal, the spiral ligament, stria vascularis and saccular macula. The cystic form of the parasite was not associated with the inflammatory response seen in the active tachyzoite form.

Conclusion
We infer that the hearing loss of toxoplasmosis is likely the result of a postnatal inflammatory response to the tachyzoite form of T gondii. Our findings have implications for the early identification and management of Toxoplasmosis.

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Role of Smac/DIABLO in TNF-α-induced Auditory Hair Cell Loss In Vitro
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Background
Second mitochondria-derived activator of caspases/Direct inhibitor of apoptosis protein (IAP)-binding protein with low PI (Smac/DIABLO) is a mitochondrial protein involved in apoptosis. Following an apoptotic stimulus, Smac is released into the cytoplasm, binds to the IAPs and releases basal inhibition of caspases to initiate apoptosis. Recently, a Smac mutation has been identified as a genetic cause of progressive non-syndromic hearing loss, DFNA64. Up to now, few studies have investigated the molecular mechanisms behind Smac activation in auditory hair cells (HC). Although Smac is known to be involved in the intrinsic pathway, studies have linked Smac expression to the extrinsic pathway through tumor necrosis factor alpha (TNFα) activity (a pro-inflammatory cytokine expressed in the cochlea in response to a variety of insults) and c-Jun N-terminal kinase (JNK).

Methods
Organ of corti (OC) explants were dissected from 3-day old rats and cultured in the following conditions: (1) No treatment(2) TNFα (1µg/ml); (3) TNFα + JNK-inhibitor (SP-600125, 10µM); (4) JNK-inhibitor alone; (5) Smac-N7 peptide (50µM); (6) Smac-N7 + Caspase-3 inhibitor (5µM); and (7) Caspase-3 inhibitor alone for either 24, 48 or 96 hrs in vitro. For HC viability, OC were fixed, and stained with FITC-phalloidin after 96 hrs in vitro and HCs with intact cuticular plates
and stereociliary bundles were counted in the basal turns. Real-time PCR was utilized to obtain Smac gene expression levels after 24 and 48 hrs in vitro. ANOVA with Tukey HSD post hoc was performed with p-value < 0.05.

Results
OC explants exposed to either TNFα or Smac-N7 displayed significantly less viable HCs in the basal turn when compared to explants in: control media; TNFα + JNK-Inhibitor; JNK-Inhibitor alone; Smac-N7 + Caspase-3 inhibitor; and Caspase-3 inhibitor alone. TNFα ototoxicity initiated time-dependent increases in Smac gene expression compared to control explant expression levels.

Conclusion
TNFα-initiated loss of auditory HCs in vitro likely involves both JNK activation and Smac expression. HC loss in OC explants treated with Smac-N7 peptide was abrogated by caspase-3 inhibitor. These results suggest a close relationship between TNFα, activated JNK, Smac, and caspase-3 activation. However, further in vitro studies involving Smac protein expression, JNK and selective caspase activation, and TUNEL assay are necessary to further delineate the role of Smac in programmed cell death in auditory HCs.

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Brain Changes that Accompany Hearing Loss
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Background
Synaptic features of the central nervous system change as a consequence of alterations in sensory input. Studies of partially deaf cats show atrophic changes in brainstem synaptic morphology and postsynaptic ultrastructure (Ryugo et al., 1998), and congenitally deaf white cats exhibit even more severe brainstem pathologies, such as synaptic hypertrophy (Ryugo et al., 1997) and a redistribution of excitatory and inhibitory synaptic terminals (Tirko et al., 2009). This study aims to confirm and expand our understanding of hearing loss as it applies to the central auditory system in mouse models of hearing loss.

Methods
Three mouse strains (aged 1-7 months) representing distinct hearing phenotypes were used: CBA/CaH, which maintains normal hearing thresholds throughout adulthood; DBA/2, which begins to exhibit high frequency hearing loss starting at ~3-4 weeks of age; and shaker-2, where homozygote mutants are congenitally deaf. Hearing thresholds were verified with auditory brainstem responses (ABRs) prior to tissue collection. Cochlear nuclei from mice of various ages and degrees of hearing loss were analyzed with light and electron microscopy. Synaptic features were labeled using glutamate transport immunohistochemistry and/or injections of neurobiotin in the auditory nerve. Spherical bushy cell (SBC) size, endbulb of Held morphology, and the distribution of axosomatic primary excitatory and/or secondary inhibitory endings were quantified.

Results
Our results show that the size of the postsynaptic SBC soma, endbulb arborization, and number of inhibitory endings onto SBCs are directly correlated to hearing sensitivity. The CBA/CaH mouse does not experience a significant elevation in hearing thresholds from 1-7 months of age, and its SBC area, endbulb arborization, and ratio of primary excitation to secondary inhibition onto SBCs remain relatively unchanged. In contrast, as hearing thresholds increase in the DBA/2 mouse, a slight decrease in all three parameters is observed. However, total auditory loss as seen in the congenitally deaf shaker-2 mouse gives rise to more drastic decreases in SBC area, endbulb arborization, and inhibition onto SBCs.

Conclusion
Preliminary data show that congenital and acquired hearing loss result in abnormal brain morphology, which can be correlated to the degree and duration of hearing loss. Based on the present results, it seems important to determine which abnormalities can be remedied by treatment methods such as sound amplification. Future studies will test the hypothesis that an amplified sound environment slows the progression of brain pathologies in comparison to that of long-term, unattended hearing loss.
Synapse Size, AMPA and NMDA Receptor Numbers, and their Distribution are Regulated in an Input-Specific Manner in the Cochlear Nucleus

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Background
Synapse size (i.e. size of the postsynaptic density, PSD) varies between synaptic contacts and often correlates with fundamental properties of synaptic transmission (Rollenhagen and Lübke, 2006; Xu-Friedman and Regehr, 2004). PSD size has been shown to be proportional to the number of ionotropic glutamate receptors (iGluRs), namely AMPA-type and NMDA-type (AMPARs and NMDARs, respectively), whose content and spatial arrangement in the synapse primarily determine synaptic quantal size in many glutamatergic synapses in the CNS (Ganeshina et al., 2004; Takumi et al., 1999). However, these anatomical, molecular and functional parameters and their correlation to electrophysiological properties of synaptic transmission in glutamatergic synapses in the binaural and monaural auditory pathways remain unknown. This information will be important for understanding normal auditory processing and for the hearing impaired.

AMPARs and NMDARs mediate excitatory transmission at auditory nerve (AN) and parallel fiber (PF) synapses in the cochlear nucleus (Raman et al., 1994; Manis and Molitor, 1996). Previously, the expression of AMPARs and NMDARs at AN and PF synapses was analyzed with preembedding immunolabeling or postembedding immunogold labeling methods (Petralia et al., 1996; Rubio and Wenthold, 1997, Rubio, 2006; Wang et al., 1998). However, neither of these techniques allows efficient detection of synaptic iGluRs. Thus, due to the technical difficulty in revealing synaptic molecules at high detection efficiency, a relationship between iGluR content and the PSD size cannot be established.

Methods
In this study we used freeze-fracture immunogold labeling (FRIL) to determine how morphological and molecular differences could relate to synaptic transmission at AN and PF synapses. FRIL allows a highly sensitive and quantitative detection of molecules of interest in the synapse (nearly 1-to-1 detection sensitivity of functional AMPARs) and provides a two dimensional (2D) landscape of the plasma membrane and receptor arrangement (Masugi-Tokita et al., 2007; Tarusawa et al., 2009).

Results
We observed substantial differences in the size of the PSD and AMPAR and NMDAR content between the AN and PF synapses on their postsynaptic targets.

Conclusion
Our results suggest that synapse size, AMPAR and NMDAR numbers and their 2D-distribution are regulated in an input-specific manner in the cochlear nucleus. These morphological and molecular properties may play important roles in shaping transmission at individual connections and normal auditory processing.

A new vertebrate-specific presynaptic protein as molecular component of the endbulb of Held

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Background
The endbulb of Held’s exceptional size has made it an important model system for the study of synaptic transmission. Furthermore, the structure of the endbulb of Held has been extensively studied to gain more insight into the process of synaptic differentiation. However, little is known about the molecular composition of endbulbs of Held. We have recently identified Mover, a vertebrate-specific 266 kDa protein, as a component of the calyx of Held. Here, we have begun to study the molecular composition of endbulbs of Held with a particular focus on vertebrate-specific presynaptic proteins, assuming that this relatively small family of proteins may confer specific features to specialized presynaptic nerve terminals. To initiate this study, we characterized the localization of Mover within the auditory pathway.

Methods
To initiate this study, we characterized the localization of Mover within the auditory pathway by using immunohistochemistry.

Results
We find abundant Mover immunoreactivity in auditory nuclei of the brainstem, e.g. the cochlear nucleus, lateral superior olive, medial nucleus of the trapezoid body, inferior colliculus. In the anteroventral cochlear nucleus (AVCN) the
immunoreactivity for Mover exhibits a punctate pattern surrounding the somas of bushy cells. The pattern of Mover is similar compared to other synaptic markers, i.e. Synapsin or Synaptophysin, but not to a glia cell marker, i.e. glial fibrillary acidic protein (GFAP). To further examine the cellular localization of Mover in presynaptic terminals in the AVCN we performed immunofluorescence staining with anti-VGLUT1, anti-VGAT and anti-Mover antibodies. At synaptic terminals ending on bushy cells we find co-localization of Mover with both VGLUT1 and VGAT.

Conclusion
Hence, we conclude that Mover is a molecular component of presynaptic endings in the anteroventral cochlear nucleus, including the endbulbs of Held.

Pulsed infrared-evoked intracellular calcium transients in cultured neonatal spiral ganglion neurons

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Background
Recent work has illustrated the feasibility of in vivo neural activation with pulsed infrared light. In the present study we analyze the response of cultured neonatal spiral ganglion cells to infrared radiation (IR), \( \lambda = 1860 \) nm, in vitro. In agreement with the previous studies, we show that pulsed IR evokes intracellular Ca\(^{2+}\) transients in the target cells.

Methods
Experiments were performed on rat spiral ganglion neurons. Neonatal animals (P2 and P3) were decapitated, and both inner ears were removed from the base of the cranium. The spiral ganglion was isolated, the cells were dissociated and plated in culture dishes coated with poly-D-lysine. After 4 days, the cells were loaded and incubated for 45 minutes with calcium-sensitive dye Fluo-4 AM dissolved in DMSO and mixed with Pluronic F-127. In a limited set of experiments, the entire organ of corti was isolated from adult rats and used to study Ca\(^{2+}\) responses. IR stimulation was achieved with a 400 \( \mu m \) optical fiber connected to a Capella laser (Lockheed Martin Aculight) operating at 1860 nm. IR was focused on the target cells under the guidance of a pilot light. Image sequences measuring fluorescence intensity differences were collected using a confocal microscope connected to a camera. The analysis of the image sequences was done with ImageJ (NIH) and Matlab (MathWorks) software.

Results
The spiral ganglion neurons responded with an increase in fluorescence synchronized in time with the pulsed IR stimulation applied. At low frequency laser pulses, a pulse-by-pulse response was observed with distinct peaks of fluorescence corresponding to intracellular calcium transients were observed. A wide range of pharmacological array was tested to identify the intracellular Ca\(^{2+}\) release site.

Conclusion
Pulsed IR consistently evoked calcium related responses in neurons. The results suggest that Pulsed IR can effectively stimulate neurons without any modifications and can find applications in neuroprostheses including in the auditory and vestibular systems.

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Stochastic properties of neurotransmitter release expand the dynamic range of endbulb synapses

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Background
The probabilistic nature of neurotransmitter release has been evident since the first recordings of synaptic potentials, but its functional effects are unknown. When the average size of synaptic potentials is near threshold, the stochastic properties of release potentially introduce noise and unreliability into postsynaptic spiking. This is important in the auditory system, where sound stimuli are encoded in the precise timing of spikes, so the loss or mistiming of spikes could reduce auditory acuity. We examined this by studying synapses formed by auditory nerve fibers onto bushy cells in the
mouse anteroventral cochlear nucleus. These synapses show significant depression, and have particularly large quanta (~100 pA) so fluctuations in the number of released vesicles could impact spiking.

Methods
All experiments were carried out using standard patch clamp methods in brain slices from CBA/CaJ mice aged P15-21 at near-physiological temperature.

Results
We measured the variability of EPSCs in voltage-clamp experiments. We then studied the impact of this variability using dynamic clamp, by comparing responses to synaptic conductances that took on the average value against those that fluctuated from trial to trial. Variability in the probability of postsynaptic spiking was almost entirely accounted for by variability in synaptic conductance, as spike threshold is highly stable. For conductances that were on average above threshold, stochastic variability reduced the probability of spiking, but for conductances that were on average below threshold, it increased spiking.

Conclusion
Our results indicated that a decreased gain allowed postsynaptic spike probability to encode more information about average EPSC amplitude, which increased the dynamic range of the synapse. Furthermore, stimuli that caused significant synaptic depression could still influence spiking because of the stochastic properties of neurotransmitter release. However, stochastic aspects of release introduced variability in the timing of postsynaptic spikes, indicating a trade-off in rate vs. time codes.

Heteromeric Purinergic P2X2/3 Channels Enhance Action Potential Firing of Developing Auditory Brainstem Neurons
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Background
In developing auditory system, purinergic signalling attunes activity in the cochlea and in the brainstem. Endogenous release of ATP contributes to action potential generation in the inner hair cells and enhances glutamate-driven firing of the cochlear nucleus (CN) bushy cells (BCs). Modulatory effects of ATP diminish with maturity, both in the periphery and in the central auditory neurons. In the brainstem, only specific cell types undergo purinergic modulation, including cochlear nucleus bushy cells, principal neurons in the medial nucleus of the trapezoid body and some neurons in the lateral superior olive. Here, activation of P2X receptor channels (P2XRs) can evoke suprathreshold membrane depolarization and action potentials by a mechanism engaging cytosolic calcium signals and subsequent activation of the protein kinase C. While the P2Rs associated with cochlear development have been largely characterized, evidence about the possible P2XR types related to maturation of central auditory neurons is still sparse.

Methods
We used in vivo/in vitro-electrophysiology and fluorometric calcium measurements to determine the exact receptor type mediating the ATP effects in the CN BCs. Experiments were performed on Mongolian gerbils, P13-16 for in vivo-, and P7-12 for slice recordings. To delineate receptor properties, we compared characteristics of native P2XR responses measured in vivo and in slices with the data obtained from HEK293 cells expressing recombinant rat P2X2R, P2X3R, or P2X2/3R.

Results
In vivo extracellular recordings in combination with iontophoretic application of AF-353, a selective antagonist of P2X2/3R and P2X3R, revealed that respective receptor subunits mediate the enhancement of spontaneous and sound evoked activity induced by the endogenous ATP release on BCs. Pharmacological characterization of responses recorded from BCs in slice preparation is consistent with P2XRs resembling mixed P2X2 and P2X3 characteristics. The desensitization profile of the currents following application of ATPγS and αβ-meATP corresponds to heteromeric P2X2/3Rs. Finally, analogous response properties were recorded in HEK293 cells expressing heteromeric P2X2/3R, but not homomeric P2X2R nor P2X3R.

Conclusion
We provide evidence that purinergic modulation of action potential firing in developing auditory brainstem is conveyed by the heteromeric P2X2/3R. Given the role of P2X2/3R in the neuronal re-organization in developing cochlea (pruning of
promiscuous hair cell-SGN connections), and the transient enhancement of spiking activity of auditory brainstem neurons shown here, it is reasonable to assume that P2X2/3R might play a more general role in development of the auditory system.

**PSD-95 Expression during Giant Synapse Formation in the Chicken Cochlear Nucleus**

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**Background**

In the cochlear nucleus, auditory nerve axons form giant synaptic terminals (Endbulbs of Held) on a specific subset of neurons only. It is yet unknown how this target-specific increase in terminal size is regulated on a molecular level during development. The “postsynaptic-density protein of 95kD” (PSD-95) is a central organizer of the postsynapse, involved in adjusting excitatory synaptic strength and size. It is also implicated in the development of the presynapse by transsynaptic interaction with neuroligins/neurexins. We thus hypothesized that PSD-95 expression should be present in the cochlear nucleus and the dynamics of expression should be temporally related to the increase in synaptic terminal size.

**Methods**

We tested this with fluorescent double immunolabeling of PSD-95 and a presynaptic marker protein (synaptic vesicle protein 2) in embryonic chicken brain cryosections from embryonic days E10 to E17. This was supported by anterograde tracing of auditory nerve axons at E15 and E16. Staining was visualized with a confocal laser-scanning microscope and analyzed with custom software.

**Results**

We found that PSD-95 was expressed at synaptic sites in all auditory brainstem nuclei of the embryonic chicken. In the mid- to high-frequency parts of nucleus magnocellularis, a maximum in the extent of PSD-95 immunosignals was evident before and at the time of maximal increase of presynaptic signal area. This was not seen in the low-frequency region of nucleus magnocellularis. Anterograde tracing corroborated that dynamics of synaptic immunosignals correlate with rapid structural changes of presynaptic terminals in the rostral regions at E15/E16.

**Conclusion**

The results support the hypothesis that PSD-95 may be involved in molecular regulation of giant synaptic development. The chicken system will allow functional studies of this protein and its binding partners in the future.
A Co-Culture of Chicken Cochlear Ganglion and Auditory Brainstem Neurons to Investigate Regulation of Endbulb Synapse Formation in Vitro
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Background
In birds, the axons of cochlear ganglion neurons project into two subdivisions of the cochlear nucleus in the auditory brainstem, the nucleus magnocellularis and the nucleus angularis. Only in nucleus magnocellularis specialized axosomatic synapses, the Endbulbs of Held, are formed. These terminals are strongly enlarged, sometimes engulfing nearly three quarters of the postsynaptic cell. The molecular basis for the determination of this giant synapse during development is still largely unknown. To address this, we established a co-culture system of cochlear ganglion (CG) and auditory brainstem neurons of the embryonic chicken.

Methods
One cell culture well was divided by an upright glass coverslip with a thickness of 130µm, creating 2 separate compartments. In one compartment auditory brainstem neurons of embryonic day 7 were seeded out. Three days later, CG neurons of embryonic day 10 were explanted into the second compartment. The coverslip was lifted after the cells had settled to allow neurite growth between compartments. Neurons were characterized via immunocytochemical staining against neurofilament and synaptic vesicle protein 2 (SV-2) at various days in vitro. SV-2 puncta size for both neuron types was measured and the CG’s neurite outgrowth onto auditory brainstem neurons was characterized.

Results
Staining against neurofilament revealed a distinct bipolar morphology of the CG neurons and a strong axon growth, reminiscent of the in vivo situation. If there were brainstem neurons present, the CG neurons showed a profound axon outgrowth into the brainstem neurons’ compartment, indicating that brainstem neurons may emit growth cues that attract CG neurons. We also could demonstrate the existence of SV-2 in the CG neurons’ axon terminals, showing that these neurons already express and transport the machinery for synapse formation as early as E10. SV-2 puncta of CG neurons appeared to be larger than those of auditory brainstem neurons and even larger when axons of CG neurons are in vicinity of auditory brainstem neurons.

Conclusion
We were able to construct a co-culture system with viable CG and auditory brainstem neurons. Our results indicate that in vitro innervation of brainstem neurons by CG neurons is possible. The differential SV-2 puncta size shows that this co-culture system is a promising tool for further investigation of molecular aspects of Endbulb-formation and developmental control of synaptic terminal size.
Structural and functional remodeling at the developing mouse calyx of Held-MNTB synapse during unilateral hearing.
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Background
Ability to locate sound is immediately impaired following a unilateral earplug but improves toward normal over several days to weeks. This suggests the reweighting of auditory cues toward the intact ear in subjects experiencing unilateral hearing deficits lead to some form of synaptic plasticity in sound localization circuits to maintain function. To examine this, we investigated the structural and functional properties at the calyx of Held-MNTB synapse in mice developing with unilateral hearing.

Methods
Experiments were performed in brainstem slices from P16-19 mice that underwent unilateral surgery to remove left middle ear ossicles at P5. Advantages to this approach are that the structural and functional properties of this synapse are well-characterized at P16-19 and because calyx terminals, which originate from globular bushy cells of the cochlear nucleus, only innervate principal cells of the contralateral MNTB, it allows for an ideal ‘in-mouse’ comparison. That being, we can quantify synaptic properties of cells located in the sound-deprived (SD) right MNTB with those located in the left MNTB (CTL) which are exposed to more auditory input.

Results
ABRs confirmed unilateral hearing deficits and could be detected at \(-10\) dB following click stimulation in normal hearing wild types (WT) and the right ear of surgical animals. Left ear ABRs had higher thresholds (65 dB) and late wave (III-V) latencies were significantly longer versus the intact right ear. Synaptic strength in CTL MNTB was enhanced based on a higher paired-pulse ratio (108%), lower probability of release (0.11) and reduced EPSC depression (25.5% of initial) in MNTB neurons following high-frequency stimulation compared to SD (respectively 95%, 0.14, 18.9%) and WT (95%, 0.17, 18.6%) synapses. Remarkably, the relationship between synaptic input and spiking properties (failure rate at 300-400 Hz) amongst the populations of MNTB neurons (WT vs SD vs CTL) were unchanged. Although failure rates were similar, SD synapses were more excitable. Calyx morphology was more complex in both the CTL and SD MNTBs compared to WTs based on the degree of branching and fenestration expected at this age. This was surprising given the classic view that the transformation from spoon to fenestrated structure is strongly driven by sound-evoked activity.

Conclusion
We characterize the structural and functional remodeling that occurs at the calyx of Held-MNTB synapse in mice developing with unilateral hearing loss as a means to maintain sound localization circuit function.

A tetanus toxin mouse model to study activity dependent processes in auditory circuitries
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Background
Neuronal activity is essential for the formation and maintenance of neuronal circuits. Previous work using cochlear ablation or gene knockouts suggested an important role of both spontaneous and acoustically driven activity for proper development and function of the central auditory system. However, these studies often suffered from ambiguity in the interpretation, as primary and secondary effects might be confounded. Furthermore, they represent irreversible processes which do not allow to challenge the plasticity of the auditory system by switching neuronal activity on and off. We therefore wished to implement a system to block neurotransmission in auditory circuits in an inducible and reversible manner.

Methods
A triple transgenic mouse model was generated to silence synaptic transmission. A Cre recombinase drives a tetanus toxin light chain (TxLC) - eGFP fusion protein in a doxycycline dependent manner. Here, we used the Egr2::Cre driver line to express TxLC-eGFP from early embryonic stages (E9.5) in rhombomeres 3 and 5 derived cells. These should result in silencing of large parts of the developing auditory brainstem such as the neurons of the AVCN, DCN, and SOC.

Results
Immunohistochemistry demonstrated expression of TxLC-eGFP in the absence of doxycyclin in the triple transgenic mouse line. Anatomical analyses of Nissl-stained sections revealed a severe reduction (41%) of the MNTB volume in adult (P25) mice. This volume reduction correlated well with a 47% decrease in cell number. These values are close to
those reported for an auditory brainstem specific Cav1.3 conditional knockout mouse, indicating an important role of excitation-transcription coupling in the developing auditory brainstem. At birth, a moderate volume reduction of 27% was already present. Currently, a similar analyses is conducted for the LSO.

**Conclusion**

Our data suggest that TxLC is expressed in the auditory brainstem and results in synaptic silencing. In addition, the anatomical results reveal that spontaneous activity is required for formation of auditory circuits. Further studies will investigate the precise mechanisms resulting in cell loss and the critical time windows for neuronal activity in auditory circuits. Finally, the approach presented here can be combined with different Cre-driver lines to block neurotransmission in any neuronal subpopulation.

### Developmental Expression of Erbb4 Is Required For Normal Cochlear Nucleus Organization

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**Background**

As the first synaptic site to receive input from the inner ear, the cochlear nucleus (CN) is a critical locus in the brain for auditory processing. Extensive literature exists on the anatomy and physiology of the CN during functional hearing, including hearing onset and decline, yet relatively less is known about the CN during its development. The laboratory is interested in elucidating molecules that contribute to the mature organization of the CN. Since the tyrosine kinase receptor Erbb4 has been shown to be a molecule critical for developmental events, including migration and differentiation, the role of Erbb4 in CN development and maturation was investigated.

**Methods**

Methods employed include in situ hybridization with digoxigenin-labeled riboprobes, general histological staining (cresyl violet, and hematoxylin and eosin) and immunohistochemistry. Analyses were performed in age-matched wild type and Erbb4 homozygous null mice.

**Results**

In situ hybridization shows that perinatal Erbb4 expression is spatiotemporally dynamic. At E16.5, expression is detected in the dorsal and ventral regions of the CN anlage. At postnatal day 0, Erbb4 mRNA is expressed in the molecular/granule cell layer and by cells within the deep dorsal and ventral regions of the CN. Examination of the mature CN in wild type and Erbb4 homozygous null mice permits assessment of the developmental impact of this gene. General histological staining shows perturbed dorsal CN organization including variable thickness of the molecular layer and altered organization of the granule cell layer. Immunohistochemical localization of vesicular glutamate transporter 1 (vGlut1; Millipore) shows vGlut1-expressing auditory nerve fiber terminals in the CN (Zhou et al., 2007) and reveals an expansion of the central core of the ventral CN in Erbb4 null mice compared to wild-type controls.

**Conclusion**

These data show that developmental expression of the receptor tyrosine kinase Erbb4 contributes to the normal anatomical organization of the CN. The change in cellular arrangement within the molecular and granule cell layers in Erbb4 (-/-) dorsal CN suggests that Erbb4 may be involved in CN neuronal migration or survival. The difference in pattern of vGlut1 immunohistochemical localization in the ventral CN between Erbb4 null and wild type mice suggests that Erbb4 may be important for the positioning of glutamatergic neurons and their afferents. Together, these anatomical changes further indicate that circuitry underlying sound localization may be critically dependent on Erbb4 expression.

### Afferent Regulation of Elongation Factor 2 in Auditory Brainstem Neurons: Cell Survival and Dendritic Reorganization

**Yuan Wang**

1. Ethan McBride

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**Background**

Afferent inputs regulate neuronal cell survival and dendritic morphology in the auditory brainstem. Using the chick nucleus magnocellularis (NM) and nucleus laminaris (NL) as models, we study the involvement of elongation factor 2 (eEF2), a translational factor that regulates protein synthesis, in afferent deprivation-induced cell death in NM and dendritic retraction in NL. Previously, we have found that afferent deprivation induced by cochlea removal leads to rapid reductions
in the level of phosphorylated eEF2 (p-eEF2) in NM neurons. The current study continues to explore the role of eEF2 activity by asking what causes observed reductions in p-eEF2 and whether these changes are associated with cell survival in NM. In addition, we examine the involvement of eEF2 in afferent regulation at a subcellular level in NL, where unilateral cochlea removal leads to selected retraction of deprived dendritic arbors but not other arbors of the same neurons with normal innervation from the other intact cochlea.

**Methods**

The chick hatchlings (P4-10) received unilateral cochlea removal and survived for up to 48 hours. Fixed sections collected from these and unoperated animals were immunostained for eEF2, p-eEF2, eEF2 kinase (EF2K, a major mechanism that phosphorylates eEF2), and phosphorylated EF2K (p-EF2K) using specific antibodies. Cell apoptosis in NM was evaluated by a dramatically decreased intensity of Nissl stain in the cytoplasm. Staining patterns in NM and NL were imaged. Average optic intensities of the staining were quantified and compared between different manipulation and survival groups.

**Results**

At 6 hours post-surgery, we found notable increases in the level of EF2K and p-EF2K immunoreactivities in the deprived NM, in parallel with dramatic decreases in phosphorylated, but not total, eEF2. In addition, we found a strong correlation of the p-eEF2 level with the intensity of Nissl staining at 12 hours in NM. In NL, we found significant decreases in the p-eEF2 level in deprived NL domain as compared to the intact domain of the same NL neurons, as early as 3 hours following unilateral cochlea removal.

**Conclusion**

Afferent inputs rapidly regulate the level of phosphorylated eEF2 in auditory neurons in NM and NL. Such regulation can be either global and associated with cell survival, or confined to specific subcellular components that undergo structural changes later. One possible mechanism underlying this regulation maybe activity-dependent eEF2 kinase signaling.
Co-release of GABA/glycine and glutamate in immature auditory brainstem from distinct vesicle populations with different release probabilities

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Background
Principal neurons of the lateral superior olive (LSO) compute interaural level differences by integrating ipsilaterally derived excitatory signals arising in the cochlear nucleus with contralaterally derived inhibitory (glycinergic) signals arising in the medial nucleus of the trapezoid body (MNTB). During the first postnatal week, glycinergic terminals in the MNTB-LSO pathway release GABA and glutamate. While GABA and glycine are known to be co-packaged, whether glutamate is co-released from the same or a different subset of vesicles has been an open question.

Methods
In acute rat brainstem slices postnatal day 4-8 (P4-8), we made whole cell voltage clamp recordings in the LSO, using reversal potentials and pharmacology to isolate different neurotransmitter responses.

Results
We examined paired-pulse ratios (PRRs) for GABA/glycine- and glutamate-mediated transmission, finding significant differences in PPR for glycine (0.66±0.03 and 0.58±0.03) and AMPA (0.79±0.02 and 0.77±0.03; p<0.001, MW-test) at 20 and 50 Hz. We next replaced calcium in the perfusate with strontium to desynchronize vesicle release and analyzed miniature synaptic events after electrical stimulation in the MNTB. In the presence of Sr++, picrotoxin, and CNQX, two populations of synaptic events could be discerned based on rise time (0.62±0.03 vs 6.59±1.3 ms at P4-5, n=12; 0.61±0.04 vs 4.96±0.25 ms at P7-8, n=12; p<0.001, Wilcoxon t) and decay time (2.17±0.1 vs 21.77±1.9 ms at p4-5, and 2.30±0.3 vs 26.1±3.3 ms at P7-8; p<001). Miniature events with fast kinetics disappeared in strychnine (10μM; n=13), whereas events with slow kinetics disappeared in D-APV (50 μM; n=11). Coefficients of variation for kinetics parameters also differed significantly for gly-R and NMDA-R events. In the presence of TTX (1μM), as with Sr++, a majority of miniature events (>98%) were fit by a single exponential. Despite fewer miniature events in TTX, we saw no differences in kinetics between evoked and spontaneous miniature synaptic events.

Conclusion
Our data are consistent with a model in which GABA/glycine and glutamate are packaged in distinct subsets of synaptic vesicles, allowing for their control by different release machinery and resulting in differing release probabilities at individual MNTB terminals.

Synaptic function in the auditory nervous system in the preterm baboon brain

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Background
Prematurity is one of the leading causes of perinatal mortality and long-term disability. Extremely premature children often suffer from developmental delays; in particular, intellectual, cognitive, academic, and language difficulties. Poor reading-speaking abilities are associated with an auditory processing disorder. Fetal auditory development is essential for early brain maturation and healthy neuronal circuitry. However, the cellular mechanisms that lead to impaired auditory processing in the developing brain are still poorly understood. We have studied developmental changes in synaptic formation and functions in the auditory nervous system in baboon brain born prematurely.

Methods
We recorded electric signals from a single synapse in preterm and full term baboon brainstem slices (125 day; 67 % gestation and 185 day gestational age controls, respectively) using whole-cell patch clamp recordings, and examined local synaptic formation in auditory brainstem using immunofluorescence microscopy.

Results
We identified multiple nuclei, the lateral and medial superior olivary nuclei (LSO, MSO) and the medial nucleus of trapezoid body (MNTB), which were prominent nuclei within the superior olivary complex under light microscope in preterm baboon brainstem slices. In preterm baboon brain, the MNTB principal neurons formed synapses with the calyx of Held terminals expressing a high density of vesicular glutamate transporters, and displayed spontaneous excitatory postsynaptic currents, which are mediated by the 2-amino-3-(3-hydroxy-5-methyl-isoxazol-4-yl) propanoic acid (AMPA) receptor.
receptors. In whole-cell patch clamp recordings, the MNTB neurons showed the burst of action potentials (~100 Hz) in response to current injections.

Conclusion
At 67% gestation, preterm baboon showed prominent neuronal circuitry and synaptic formation in the auditory brainstem, thus the functional maturation of synapses during late gestation period is critical for the development of auditory processing.

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A biophysical cable model for MSO neurons as generators of the neurophonic
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Background
The medial superior olive (MSO) is an early stage of binaural processing and sound localization in mammals. In vivo recordings of single cell activity in the MSO are rare due to the presence of strong sound-evoked local field potentials (LFPs) in the brain stem. These LFPs, known as the auditory neurophonic, are believed to reflect post-synaptic current flow in MSO neurons and are a more accessible signal of in vivo neural activity. To date, however, the exact generators of the LFPs remain unclear, and there has been no quantitative theory that links the neurophonic to activity of individual MSO neurons.

Methods
We use biophysically-based cable models of MSO bipolar neurons to predict how single cell neural dynamics can generate the experimentally observed population-level neurophonic (McLaughlin et al 2010). Passive cable theory (Rall 1977, e.g.) provides initial insights which are complemented by simulations of a model that includes voltage-gated currents (Mathews et al 2010). Motivated by anatomical and neurophysiological observations we make two simplifying, yet plausible, assumptions: 1) MSO neurons are identical bipolar cells arrayed with spatial symmetry; 2) afferent inputs to a subpopulation of MSO cells are in synchrony with each other. With these assumptions, we formulate a one-dimensional model of the population-summed neurophonic and relate simulation results to in vivo data.

Results
We find that post-synaptic dendritic current flow in the MSO neuron models can generate extracellular potentials that qualitatively resemble LFPs evoked by monaural tones (see figure). Our simulations make some non-intuitive predictions for responses to binaural beat stimuli. For instance, the spatial spread of LFPs depends on the degree of coincidence of bilateral inputs with coincident inputs generating fields that are most localized near the model MSO neurons. In ongoing work, we use these predictions to help interpret experimental data.

Conclusion
Our modeling provides a biophysically-based foundation to assess how cellular and synaptic properties generate the neurophonic. We have shown that, for monaural stimuli, the bipolar morphology of MSO neurons can create dipole-like LFPs with a half cycle phase shift between sink and source. This confirms and extends earlier conceptualizations of the neurophonic (Galambos et al 1959, McLaughlin et al 2010). These findings contrast with a study of the neurophonic in barn owls which suggested that pre-synaptic currents are the dominant generators of brain stem LFPs in that system (Kuokkanen et al 2010).
In Vivo Whole-Cell Recordings from Principal Neurons of the Medial Superior Olive

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Background

In vivo intracellular data from medial superior olive (MSO) neurons have not been reported. Such data are essential to understand how sensitivity to interaural time differences (ITD) is generated in these cells. Its absence has led to controversies, for example regarding the origin of internal delays and the nature of the coincidence process.

Methods

We performed in vivo whole-cell current clamp recordings in the superior olivary complex (SOC) of young gerbils under general anesthesia. Patch clamp electrodes were filled with an internal solution containing biocytin. Neurons were characterized with threshold tuning curves, monaural and binaural sinusoidal tones, binaural beats and broadband noise. We separated postsynaptic potentials (PSPs) and spikes in the voltage signal using criteria of amplitude and repolarization speed. At the end of the experiment the animal was perfused and the brainstem sectioned, so that biocytin-filled cells could be retrieved using routine histological methods.

Results

We obtained in vivo intracellular voltage responses for a variety of SOC neurons. Sensitivity to ITD as well as histology (cf. Figure) confirmed several of them to be MSO principal neurons. Analysis of the spontaneous activity showed remarkably high and continuous subthreshold activity. Driven activity, both to monaural and binaural stimulation, consisted of discrete excitatory events rather than a stimulus-like analogue waveform. Inhibitory PSPs were not obviously apparent as discrete events, either in the spontaneous signal or in the acoustically driven signal. The rise time and duration of subthreshold excitatory PSPs were close to the values previously reported for MSO neurons in brainstem slices. By comparing the response to ipsilateral and contralateral sounds, we found that rise times of subthreshold events triggered from opposing ears were not consistently different. The precision of phase locking (the vector strength of spikes) was higher than that of subthreshold events. The relation between characteristic frequency and histological location shows that the high-frequency representation is surprisingly limited.

Conclusion

In conclusion, we made the first in vivo whole-cell recordings from MSO neurons. While our results do not preclude a role for inhibition, they suggest that ITD sensitivity is determined primarily by the relative timing of discrete, similarly shaped EPSPs.

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Activity dependent duration of inhibition in an echo processing circuit

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Background
To localize primary sound sources in reverberant environments, the auditory system has to suppress the directional information of echoes. An elementary step of this process seems to be implemented in the circuitry of the dorsal nucleus of the lateral lemniscus (DNLL). The DNLL generates an exceptionally long lasting GABAergic inhibition that suppresses responses to trailing sounds for tens of milliseconds in the contralateral DNLL termed persistent inhibition (PI). It has been suggested that this PI underlies the suppression of directional information during echoes. However, in vitro single GABAergic IPSC decay with a time constant of \(\sim 4\) ms, being too fast to explain the PI. Data from juvenile gerbils suggest that the appropriate inhibitory time course might be adjusted in an activity dependent manner. In this study we describe the cellular basis of the PI and its dependence on activity.

Methods
The cellular basis of PI was analyzed using in vitro patch clamp recordings in acute brain slices from P30-35 Mongolian Gerbils at near physiological temperatures. To do so recordings of pharmacologically isolated, fiber stimulated GABAergic responses as well as current and conductance injections into postsynaptic DNLL neurons were carried out.

Results
The GABAergic IPSC decay time course slowed with increasing stimulation frequencies, strength and pulses number indicating an activity dependent process. Pharmacological dissection of the underlying synaptic mechanisms showed that transmitter spill over and asynchronous release mediate the increase of GABAergic decay time constants. The factors that influence the conversion of this inhibitory conductance into effective IPSPs were analyzed. GABAergic IPSPs are strongly hyperpolarizing, as the chloride reversal potential is \(\sim -90\) mV. The membrane properties between the resting potential and the reversal potential indicate that synaptic GABA conductances are integrated passively. A steady state approximation based on these findings shows that larger synaptic conductances cause slower IPSPs. The injection of synaptic conductances shows that these membrane properties transform the activity dependent slowing of IPSCs into a long lasting hyperpolarization. Finally, this hyperpolarization is indeed sufficient to suppress action potentials for tens of milliseconds.

Conclusion
Taken together, our data indicate that activity-dependent, hyperpolarizing inhibition can mimic the persistent inhibition in vivo underlying the suppression of space information during echo perception.
Systematic variation of conduction velocity achieves isochronic inputs
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Background
Proper function of brain circuitry relies on the exact timing of signal propagation. In the avian brainstem, the circuit responsible for sound segregation consists of internal axonal delay lines innervating an array of coincidence detector neurons that encode interaural time differences (ITDs). Information from the ears gets transferred to nucleus magnocellularis (NM) neurons on each side of the brainstem and individual NM axons project to the dorsal dendritic region of the ipsilateral nucleus laminaris (NL) and to the ventral region of the contralateral NL, resulting in NL neurons receiving segregated input from both ears. These pathways are thought to represent a circuit similar to the Jeffress Model of sound localization. This model assumes equivalent delays from the two ears when a stimulus originates from straight ahead, leading to ITDs of less than 10 µs. To accomplish this, action potential (AP) speed and travel distance have to be timed precisely - in the microsecond range - to ensure coincident arrival of information at individual detector neurons in NL.

We previously reported that internode distance and axon diameter of NM neurons vary systematically within individual axons to compensate for axonal length differences, suggesting that conduction velocities of specific axon segments are precisely regulated; shorter axon segments should propagate APs slower than longer axon segments.

Methods
In the current study, conduction velocities in different NM axon segments were measured. To this end, we recorded from single NM neurons in whole-cell current-clamp mode and determined the difference in the travel time of antidromic APs, elicited at two different locations along the axon. The axons were visualized with biocytin, and with the axon segment length between the stimulation sites conduction times were determined.

Results
Our results show that longer contralateral axons projecting across the midline conduct APs faster than the shorter axons in the ipsilateral loop. Conduction velocities calculated for physiological temperatures and axon length measurements show that conduction times of the ipsilateral and the contralateral NM axon branch are almost equal, leading to coincident arrival of inputs at NL when ITD is 0.

Conclusion
These combined anatomical and functional properties lead to the arrival of appropriately timed binaural inputs to coincidence detector neurons. Conduction velocity is systematically regulated within two different branches of the same axon and the parameters responsible for conduction velocity of NM axons must be regulated precisely during development to achieve coincidence detection.

A Model MSO Neuron Approaches the Measured Frequency Range of Fine-Structure ITD Sensitivity
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Background
Sound from an acoustic source located to the side arrives first at the near ear, producing an interaural time difference (ITD). Sensitivity to ITD is derived in the left and right medial superior olive (MSO) of the auditory brainstem. MSO neurons receive bilateral excitation via the anteroventral cochlear nuclei (AVCN), and inhibition via the medial and lateral nuclei of the trapezoid body (MNTB and LNTB, contralaterally and ipsilaterally driven, respectively). For pure-tone acoustic inputs, in-vivo measurements in cat MSO have focused on frequencies from 150 to 1000 Hz, and recently in gerbil MSO from 800 to 2000 Hz. Fine-structure ITD sensitivity to 2 kHz is reported in both species, and effectively modeled above 800 Hz using bilateral excitation (EE) and frequency-dependent cochlear delays. This study models neural mechanisms underlying ITD sensitivity within an MSO neuron incorporating realistic action potential propagation and an accurate number of synaptic inputs.

Methods
Existing ion-channel dynamics equations from gerbil MSO neurons were incorporated into a Hodgkin-Huxley model MSO neuron comprising two dendrites (contralateral and ipsilateral), soma, and axon. Four excitatory synapses were applied to each dendrite, zero to eight inhibitory synapses to the soma. Synaptic strength and speed were adjusted. Each synapse was driven by a point process with discharge patterns reflecting an AVCN bushy cell, or TB neuron, stimulated at
characteristic frequency at 60 dB SPL. Active channel mechanisms were emphasized by minimizing passive leakage currents at resting potential, which was slightly lower in the axon than in the dendrites and soma.

Results
Large action potentials were limited to the model axon with minimal back-propagation to the soma. At 150 Hz, the model MSO neuron with bilateral excitation and inhibition (EEII) produced a broadly tuned rate-ITD function ranging from 0 to 120 spikes/s and monaural responses around 50 spikes/s, matching measurements in cat MSO. At 1600 Hz, using faster synapses the EEII model MSO neuron matched rate-ITD functions from gerbil MSO (ranging from 10 to 100 spikes/s), but monaural responses were weaker than in gerbil. From 300 to 1250 Hz, bilateral excitation and contralateral inhibition captured ITD responses in cat and gerbil MSO.

Conclusion
The model MSO neuron matched rate-ITD functions for pure-tones approaching the frequency range of fine-structure ITD sensitivity. Ion-channel mechanisms restricted back-propagation of action potentials, and supported ITD sensitivity for an accurately small number of synaptic inputs.
### The Influence of Waveform Envelope Shape on the Rate-ITD Functions of Neurons in the Inferior Colliculus

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**Background**

It is possible to design envelope waveforms with each segment - pause, attack, sustain, and decay – assigned an independent duration. Klein-Hennig et al. (2011, J. Acoust. Soc. Am. 129 p. 3856-3872) conducted psychoacoustic experiments utilising these envelope waveforms to determine the threshold interaural time differences (ITDs) for different waveform envelope shapes and revealed that a short duration attack segment preceded by at least 4ms of pause is optimal for discrimination performance, whilst sustain and decay segment durations were of only minor influence.

**Methods**

Using tungsten microelectrodes, we recorded rate-vs.-ITD functions of single neurons in the inferior colliculus (IC) of anaesthetised guinea pigs. For each neuron, carrier frequencies corresponded to the neuron's characteristic frequency (CF). Responses for 3 envelope shapes were analysed: (1) short attack-long decay (1.5ms-15ms), (2) long attack-short decay (15ms-1.5ms), and (3) a short attack-short decay (1.5ms-1.5ms) pseudo square-wave envelope. The current study assesses how each envelope waveform segment influences the rate-ITD function of IC neurons for each envelope shape.

**Results**

Rate-ITD-functions for the short-long stimuli (1) were better modulated than those to the long-short (2), indicating improved ITD discriminability in the presence of a waveform envelope with a short attack segment duration. Each neuron for most neurons, the shape of the rate-ITD functions, and therefore ITD sensitivity, for the various envelopes differed considerably: a wide range of neural ITD sensitivity to different is unique in the shape of its rate-ITD-functions for each envelope shapes was observed. The neural ITD discrimination threshold is here defined as the smallest significant difference in response rate to any reference ITD. The responses of the first 28 neurons to be measured revealed 21/28 to have a significant (d'≥2) ITD discrimination threshold for the short-long stimulus compared to 8/28 for the long-short stimulus. Average thresholds for the neurons in the top quartile of this subset were 167 µs for the short-long and 1000 µs for the long-short stimulus.

**Conclusion**

The data indicate the importance of fast-attack envelopes in providing for high-resolution ITD sensitivity in high-CF neurons. The physiological data show strong correspondence with human performance in envelope-ITD discrimination tasks.

### GABA<sub>B</sub> receptor mediated adaptation in medial superior olive neurons

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**Background**

It is well established that human listeners adapt in their judgement of the lateral position of low-frequency sound sources depending on the nature, timing and location of preceding sound stimulation. Recently it has been shown that interaural level difference processing in the lateral superior olive (LSO), which is important for localizing high-frequency sounds, is subject to binaural adaptations mediated by GABA<sub>B</sub> receptors (Magnusson et al. 2008). The location of low-frequency sounds is first encoded in the medial superior olive (MSO) that precisely detects interaural differences in the arrival time of a sound at the two ears (ITDs). GABA<sub>B</sub> receptors are expressed in the MSO, but in contrast to LSO neurons it is unclear whether MSO neurons also adapt their binaural sensitivity.

**Methods**

Here, we performed human psychophysical experiments and in vivo extracellular recordings of single MSO neurons in adult Mongolian gerbils to investigate adaptation in ITD-based sound localization using the same stimulus paradigm in both approaches. Moreover, we applied pharmacological manipulations in vivo and in vitro to determine the synaptic mechanism underlying neuronal adaptation.

**Results**

Our human psychophysical and in vivo electrophysiological studies show that adapter tone pips change human perception of the location of test tones and spike rates of gerbil MSO neurons, respectively. Specifically, we find that the spike rates...
during the test tones were negatively correlated with the spiking activity of individual MSO neurons to the adapter, suggesting an activity-dependent adaptation. Importantly, a similar reduction in spiking activity as found during the tone-evoked adaptation could be elicited by iontophoretically applying GABA without presenting adapter sounds. Moreover, the overall spike rate decreased strongly during application of the GABA<sub>B</sub> receptor agonist baclofen, and conversely, antagonizing GABA<sub>B</sub> receptors with CGP 55845 increased spiking activity. In line with this, in vitro patch-clamp studies revealed that GABA<sub>B</sub> receptors modulate both excitatory and inhibitory inputs to the MSO in a dose-dependent manner.

**Conclusion**
Taken together, our results suggest that MSO neurons adapt their spike rate according to their preceding spiking activity through a GABA<sub>B</sub> receptor mediated feedback loop. This feedback loop might serve as a dynamic adjustment during changing environments.

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NMDA-dependent enhancement of rate coding in the auditory brainstem.**
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**Background**
Low frequency neurons in the dorsal nucleus of the lateral Lemniscus (DNLL) are sensitivity to interaural time differences. They directly inherit this sensitivity from neurons of the superior olivary complex. Coincidence detector neurons in the superior olivary complex translate the binaural temporal code into a rate code. A major role of DNLL neurons at the second stage of this binaural pathway is suggested to be enhancing the rate code generated at the site of coincidence detection (Pecka et al. 2010). This enhancement is due to a reduction in the variability of neuronal responses in the DNLL compared to the superior olivary complex. However the cellular mechanisms of this enhancement are unclear. Interestingly, recent in vitro data from our lab showed that the synaptic NMDA currents amplify the postsynaptic activity in adult DNLL neurons (Porres et al. 2011, Ammer et al. 2012).

**Methods**
In this study we now linked the in vitro results to the in vivo measured enhancement investigating the role of NMDA currents on the neuronal activity of DNLL neurons in adult Mongolian gerbils using a combined in vivo and in vitro approach. While pharmacological blocking the NMDA current we measured the contribution of NMDA receptor mediated activation on the binaural responses in terms of rate, time, and variability in vivo.

**Results**
In vivo the neuronal activity in adult gerbils is significantly reduced after blocking the NMDA current. Besides the onset of the response to pure tones is strongly reduced by applying NMDA antagonists. This reduction of the ongoing component resulted also in shorter responses to tone stimulation. Furthermore the fano factor, a measure of response variability, significantly increases after blocking the NMDA current, which implies a higher variability in the neuronal response after blocking. The NMDA receptor induced enhancing of the rate code directly enhanced the neuronal coding of interaural time differences. This NMDA dependency is in line with in vitro findings that show a reduction in the number of spikes in response to fibre stimulation in a frequency dependent manner.

**Conclusion**
Taken together a NMDA sensitive component amplifies neuronal activity in the DNLL. This supports a transition from a predominantly temporal to rate-code information at this second stage of the binaural pathway.

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Axonal Organization of Inhibitory Inputs to Binaural Brainstem Circuit**
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**Background**
A fundamental question in neuroscience is how connections between neurons underlie information processing. In the chicken auditory brainstem, excitatory and inhibitory inputs to the nucleus laminaris (NL) are thought to be organized in a way that optimizes the computation of interaural time differences (ITDs), a binaural cue for encoding the location of a sound source. Neurons in NL are bipolar, with dorsal and ventral dendritic lamina receiving excitatory input from separate
ears. Excitatory axons innervate narrow bands across NL to construct a precise tonotopic map. Orthogonal to its tonotopic map, NL is arranged by preferred ITD such that neighboring neurons have adjacent receptive fields along the horizontal plane. Computational and physiological studies demonstrated that the sensitivity of NL neurons to ITDs is modulated by an inhibitory feedback circuit via the superior olivary nucleus (SON). To better understand the structural arrangement of inhibitory inputs that provide this modulation, we determined the pattern of axonal arborizations of individual SON axonal arborizations innervating NL.

**Methods**

We performed in vitro intracellular dye filling to label individual SON neurons and their terminal axonal arborizations in NL in embryonic day 18-21 chickens. Immunolabeling for GABAergic synapse components was performed on this tissue to ascertain the spatial distribution of synaptic contacts of single SON axons. Images were acquired using fluorescence microscopy and analyzed using FIJI and Matlab software. Measurements of arborizations (i.e. length, area, shape and location) and synaptic patterning (i.e. density and location) were compared between tonotopic regions and the two dendritic laminae of NL.

**Results**

The arborization patterns of SON axons were largely variable. While some axons targeted a specific dendritic lamina of NL (similar to excitatory inputs), others innervated both laminae relatively equally. In addition, some SON axons innervated a specific frequency region of NL while others extended across almost the entire tonotopic axis. SON axons in the later group did not innervate all frequency regions uniformly, but instead showed a high density of branches and synapses in a confined region and a significantly lower density of branches and synapse in surrounding frequency regions.

**Conclusion**

In contrast to the highly stereotyped and precisely targeted excitatory inputs to NL, the spatial patterns of inhibitory inputs from SON to NL are diverse. Further analyses will quantify the patterns of axon arborizations. This information will provide a more thorough understanding of how inhibitory input enhances frequency mapping and spatial segregation in ITD pathways of the brainstem.
Frequency Region Differences in Outward Currents of the Nucleus Laminaris in Chicken.
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Background
In order to determine the location of a sound in horizontal space, specialized neurons in the auditory system compare arrival times of inputs (interaural time differences; ITD) from both ears. In chick, neurons in the nucleus laminaris (NL) compute ITD along a tonotopic axis. In vitro data suggests middle and high characteristic frequency (CF) neurons encode ITD more accurately than low CF neurons. It has been proposed that the presence of strong Kv currents partially explains this observation. The purpose of this work was to further study Kv currents in NL neurons across the tonotopic axis in order to better understand their role in ITD coding.

Methods
Brainstem slices (275-300 µm) were prepared from chick embryos (E18). An Axopatch 200B amplifier was used to perform whole-cell voltage-clamp experiments. Recordings were made at room temperature, in the presence of TTX (1 µM), and at a holding potential of -60 mV. Leak subtraction was implemented using a linear regression of current measured between -100 and -85 mV.

Results
Total Kv-currents in high CF neurons (at 0 mV, 12.0 ± 2.4 nA) were larger than in low CF neurons (at 0 mV, 7.1 ± 1.17 nA; p = 0.03). After accounting for size differences between cells, data confirmed that high CF neurons (at 0 mV, 0.34 ± 0.06 nA/pF) possess more total-Kv current density than low CF neurons (at 0 mV, 0.15 ± 0.05 nA/pF; p < 0.01). Furthermore, the V1/2 of total Kv current was substantially more hyperpolarized in middle (-16.9 ± 3.7 mV) and high CF (-15.4 ± 3.5 mV) than low CF neurons (1.2 ± 4.3 mV; p < 0.001). High CF neurons also possessed more Kv current at rest. Despite variations of voltage-gated sodium and Kv channels along the tonotopic axis, the two currents did not co-vary along the frequency axis. Finally, a small inactivating component of Kv current was present in middle and high but rarely in low CF neurons.

Conclusion
The results of this study indicate that high CF neurons in the NL possess stronger and more active Kv current than low CF neurons. Interestingly, there were few significant differences between middle CF neurons and low or high CF neurons. More work is needed to examine Kv currents in adult chicken and address how Kv currents improve ITD coding across different frequency regions.

Group II mGluRs Induce LTD of GABAergic Transmission in Avian Auditory Brainstem Neurons
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Background
Since its discovery, chemical long-term depression (LTD), a persistent reduction in synaptic efficiency induced by agonists for transmitter receptors such as metabotropic glutamate receptors (mGluRs), has been extensively studied at excitatory glutamatergic synapses. However, group II mGluR (mGluR II)-induced LTD at inhibitory synapses has not been reported. We previously found that mGluR II agonist DCG-IV suppressed GABAergic IPSCs of chicken cochlear nucleus magnocellularis (NM) neurons, and recovery after washout of the agonist (~20 min) was incomplete. We hypothesized that mGluR II induced LTD of GABAergic transmission in NM neurons.

Methods
Brainstem slices (300 µm in thickness) were prepared from chick embryos (E17-20). Whole-cell voltage-clamp experiments were performed with an AxoPatch 200B amplifier, at a holding potential of -70 mV, at 34-36 °C. IPSCs were recorded (once every 30 s, for > 1 h) in the presence of antagonist for AMPA receptors.

Results
DCG-IV (4 µM) induced a strong initial inhibition followed by LTD of IPSCs, which was blocked by mGluR II antagonist LY341495. DCG-IV application without synaptic stimulation of the GABAergic afferents failed to induce LTD, indicating that presynaptic spike activity was required to form the LTD, and more importantly, DCG-IV was washed out rapidly. Analyses of the failure rate and coefficient variance (CV) of IPSCs revealed presynaptic actions of mGluR II. NMDAR antagonist APV did not change the LTD, indicating that the LTD is NMDAR independent. Blocking adenylyl cyclase and
PKA by pre-incubating slices for 1 h in ACSF containing SQ22536 and KT5720, respectively, eliminated the LTD. Surprisingly, blocking mGluR II increased the frequency without affecting the amplitude of mIPSCs, suggesting that ambient glutamate activated mGluR II. A low frequency stimulation (LFS, 1 Hz, for 15 min) protocol that activated both the glutamatergic and GABAergic pathways to NM induced LTD, which was blocked by LY341495, confirming chemical LTD induced by mGluR II activated by synaptically released glutamate and excluding electrical LTD induced by LFS of the GABAergic afferents.

Conclusion
Our finding of mGluR II-induced LTD of GABAergic synapses represents a novel form of long-term plasticity in the central auditory system, and provides evidence supporting the model we previously proposed in which tonic activity of mGluRs regulates the GABAergic transmission in NM, enhancing temporal information processing of sounds. Supported by NIH NIDCD Grant DC008984 to YL.

### Action potential initiation in MSO principal neurons examined with light-regulated voltage-gated ion channel blockers

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**Background**
The principal neurons of medial superior olive (MSO) compute microsecond temporal differences in arrival time of sounds to the two ears. The axon initial segment (AIS) is thought to be the site of action potential initiation in MSO neurons, but despite its crucial position, the technical difficulty in isolating the effects of voltage-gated ion channels in the AIS from those of the soma and dendrites has precluded a clear understanding of how AIS properties influence the coding of auditory information.

**Methods**
We examined action potential initiation in gerbil brainstem slices containing the MSO at 35°C. To investigate the specific functional roles played by the AIS, we combined whole-cell patch recordings and confocal imaging with recently developed photoswitches (e.g. QAQ) which provide a wavelength-dependent block of voltage-gated Na, K, and some Ca channels at specific subcellular locations. Both QAQ (200 µM) and Alexa 568 (50 µM) were applied through the pipette via diffusion and allowed to equilibrate in MSO neurons for >10 min., and neuron morphology was visualized. Simulated EPSCs were delivered through the recording electrode both during field illumination at 380 nm and after ion channels were focally blocked by scanning a 488 nm laser over regions of interest. These effects were immediately and fully reversible upon return to 380 nm illumination.

**Results**
Using intracellular QAQ, we compared the effects of ion channel blockade at either the AIS, soma, or entire cell. In slices from P13~14 gerbils, 488 nm illumination of the axon up to 50 µm from the soma induced either a complete failure in action potential initiation or an increase in voltage threshold (-47 to -39 mV, p=0.0002, n=6). By contrast, somatic block provided a far smaller increase in threshold (-47 to -45 mV, p=0.015; n=6). Similar results were obtained in MSO neurons from gerbils >P18. These results are consistent with the primary effects of QAQ on the voltage-gated Na channels underlying the action potential.

**Conclusion**
We find that action potential threshold is established primarily by Na channels in the AIS, with surprisingly little contribution from somatodendritic Na channels. Thus spike initiation in the AIS appears electrically isolated from the influence of Na channels in the soma and dendrites.
**Stimulus Specific Adaptation in the Auditory Thalamus of Awake Rats**

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**Background**

Novelty detection by single neurons in the mammalian auditory system has been suggested to be a real-time filtering or gating mechanism of acoustic information. Selective adaptation of a neuron’s response to repetitive/high-probability stimuli, but not rare/low-probability stimuli in a given domain is known as stimulus specific adaptation (SSA). Quantification of SSA using specific indices and stimulus protocols has allowed for the quantification of a single neurons’ ability to respond selectively to novel stimuli. As defined by these metrics, SSA occurs in the inferior colliculus, medial geniculate body (MGB) and auditory cortex of anesthetized rodents. However, only auditory cortex has been examined and found to display SSA in awake animals.

**Methods**

Individually-advanceable tetrode microdrives were implanted to record single unit responses in the auditory thalamus (MGB) of awake Fischer Brown Norway rats. To determine the presence of SSA in MGB of awake rats, two stimulus paradigms were used to evoke SSA: the oddball paradigm and isointensity random and non-random frequency function pairs.

**Results**

Single units in the MGB of awake Fischer Brown Norway rats display SSA in response to two different paradigms. SSA was stronger in the non-lemniscal regions of the MGB and was most dramatic at lower intensities where 27 of 57 (47%) young adult single units displayed SSA. However, when SSA level, intensity and time course were compared to SSA responses of aged MGB single units, there were no significant age-related differences.

**Conclusion**

These data provide a description of SSA in the MGB of awake rats using two different stimulus SSA-inducing paradigms. Findings indicate that SSA is not dependent on arousal level nor the anesthetized state, but is a common response in the MGB of awake rats. SSA did not appear to be overtly altered in the aged auditory thalamus of awake rats. Determining the relevance of SSA to acoustic scene coding in a whole animal, what region in the auditory neuraxis and what neurotransmitters, receptors and membrane conductances are responsible for the generation of SSA remain as primary goals in the study of SSA.

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**Stimulus-Specific Adaptation in Real and Model Neurons**

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**Background**

Stimulus-specific adaptation (SSA) is the reduction in the responses to a repeated stimulus that is not, or only partially, generalized to other stimuli. SSA is extremely sensitive, showing frequency hyperacuity, and being induced by frozen tokens of white noise ‘tone clouds’. However, the mechanisms underlying SSA are still unclear. Along the non-lemniscal pathway, SSA is found as early as the inferior colliculus (IC), but along the lemniscal pathway, SSA first appears in primary auditory cortex (A1). Two different mechanisms may therefore underlie the generation of SSA in the lemniscal and the non-lemniscal pathway.

**Methods**

We studied two possible models for the generation of SSA. The first model is feed-forward, based on adaptation of excitation in narrow frequency channels that impinge on a single output neuron. The second is a recurrent network model, based on the tendency of networks with depressing synapses to generate ‘population spikes’. Population spikes are more likely to occur for rare than for common stimuli, giving rise to SSA. We generated predictions for responses to oddball and control sequences from both models. We compared the responses predicted by these models to responses recorded extracellularly from the IC, and intracellularly and extracellularly from auditory cortex of halothane-anesthetized rats.

**Results**

The feed-forward model predicts responses to a deviant tone to be smaller than responses to the same tone occurring with equal probability among many others (“deviant among many standards”). In IC, the feed-forward model fits nicely the data: neurons showing strong SSA are widely-tuned, as expected from the integration of many narrowly tuned inputs; and
the responses to deviants in oddball sequences are indeed smaller than the responses to deviants among many standards. In contrast, in cortex, the predictions of the feed-forward model fail to a large extent: responses to deviants are larger than predicted, and are essentially equivalent to the responses to deviants among many standards, both for membrane potential responses and for spiking responses. The network model, on the other hand, reproduces the cortical results.

Conclusion
SSA in IC and in cortex seem to be due to different mechanisms. While in IC, widely-tuned neurons show SSA that is compatible with feed-forward adaptation of excitation in narrow frequency channels, A1 neurons do not conform to this model, but seem to be reasonably well described by a model that relies on network dynamics for the generation of SSA.

A Functional Near-Infrared Spectroscopy Study (fNIRS) of the McGurk Effect in Normal-Hearing Adults

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Background
Human speech perception is multimodal in nature, involving cross-modal binding of auditory and visual input. The McGurk effect can be elicited when incongruent audio-visual information, such as visual “ga” and audio “ba”, are integrated and result in a novel fused percept that differs from both the audio and visual percept, such as “da”. Functional MRI studies have correlated greater cortical responses in the auditory cortex with perception of the McGurk effect. In this study we used fNIRS to measure hemodynamic responses in the auditory cortex of normal-hearing adults to a stimulus designed to elicit the McGurk effect (McGurk stimuli) and stimuli that does not elicit the McGurk effect (non-McGurk stimuli).

Methods
Behavioural tests were performed in order to establish the efficacy of the McGurk stimuli. Adult normal-hearing participants (n=18, female n=12) were presented with the McGurk stimuli (audio “ba” with visual “ga”) and were asked to report the syllable that they perceived in an open-choice response task. In a separate session, we assessed fNIRS responses to McGurk and non-McGurk stimuli. Non-McGurk stimuli included audio alone “ba”, visual alone “ga”, and audio “ga” with visual “ba”.
During the fNIRS experiment, hemodynamic responses were measured using 3x3 arrays of optodes (Hitachi-ETG4000 system) placed bilaterally over the temporal lobes. All stimuli were presented five times in a pseudorandom order and interleaved with a baseline condition.

Results
The behavioural response task revealed that in 78% of subjects the McGurk stimuli elicited a novel fused percept that neither matched the audio nor the visual input. Fewer subjects reported a percept that matched the auditory component of the McGurk stimulus (17%) or the visual component (5%). Thus far we have completed fNIRS measurements in ten participants. Ongoing analysis will compare hemodynamic responses in the auditory cortex to the McGurk and non-McGurk stimuli.

Conclusion
The 78% rate of susceptibility to the McGurk effect is consistent with previously reported rates. Therefore our findings suggest that the present McGurk stimulus is an effective stimulus for eliciting the McGurk Effect in normal-hearing adults. Our future research aims to acquire fNIRS data in a cohort of cochlear implanted adult subjects. Accordingly, we aim to compare hemodynamic responses to McGurk stimuli in the auditory cortex of normal-hearing and cochlear implanted adult populations.

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Plastic changes of the auditory core fields after behavioral conditioning to a natural sound
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Background
Reversing in time is sometimes used to elucidate whether neuron's responsiveness to a given complex sound is specific or not. In electrophysiological recordings made from auditory cortex (AC) of untrained animals, some neurons show dissociated (different) and others show associated (similar) responses when pairs of natural sound and its time-reversed version (REV) are used as stimuli. However, it is not known whether the same response properties are still preserved in conditioned cortex. There may be the possibility that the conditioned stimulus would change the neuronal network and that the changed network might facilitate or suppress activation evoked by its REV. We compared the activation properties evoked by such a stimulus pair between naive and fully conditioned AC.

Methods
Guinea pigs (Hartley, 300 to 500g) were operant-conditioned to a digitized footstep sound (FSS) with food as reinforcer, resulting in 2 weeks in the induction of distinctive response behaviors in more than 95% trials. These included quick head swaying and/or circling locomotion initiated soon after the onset of conditioning sound. AC including core fields (CORE) of both untrained naive and trained groups was subject to voltage-sensitive dye optical imaging under ketamine anesthesia.

Results
Animals conditioned to the FSS did not respond behaviorally to its REV (rFSS) in most trials. Naive animals did not respond behaviorally to either stimuli at all. Optically-measured neuronal activity was evoked across the CORE to either stimulus type whether animals had been trained or untrained, but its extent and strength were varied between the stimulus types and also between the training experience. A remarkable contrast in response pattern to FSS and rFSS was found in the conditioned cortex, with a strong activation to FSS and a week activation to rFSS. On the contrary, in the naive animals, the strong activation was evoked by the FSS, like in conditioned animals, but the activation evoked by rFSS was relatively strong, being generally greater than that evoked in the conditioned cortex.

Conclusion
Although neuronal mechanisms underlying this asymmetry in the response contrast to the different stimuli between the conditioned and non-conditioned cortex are currently unknown. One plausible explanation would be that a potential change in neuronal network is induced by the sound used for conditioning (FSS) and that this plastic change might not permit the sound with the opposite temporal structure to activate the cortex or subcortical stations. Supported by KAKENHI to H.O. (no.22500368).
Task-dependent activations of human auditory cortex during pitch and spatial tasks

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Background

It is generally accepted that spatial and nonspatial auditory information is processed in separate streams in human auditory cortex. However, our previous studies have shown that activations during spatial discrimination and spatial memory tasks are highly similar to those observed during pitch discrimination and pitch memory tasks (Rinne et al, 2009, 2012). In the present study, we systematically compared fMRI activations during discrimination and memory tasks performed on sounds varying in both pitch and location. In different conditions, subjects performed the tasks on sounds in which both features varied or only the task-relevant feature varied and the task-irrelevant feature was absent (not salient). To observe pure task effects, subjects were also instructed to perform the discrimination tasks on sounds in which both features were absent. In addition, the different stimulus conditions were presented during a visual task. We expected that this paradigm would reveal task-dependent and stimulus-dependent activations associated with spatial and nonspatial processing.

Methods

In different blocks, subjects (N = 22) performed pitch/spatial discrimination, pitch/spatial memory or visual tasks. Auditory stimuli were noise bursts (duration 200 ms, mean rate 0.9 s) consisting of two 90-ms parts separated by a 20-ms gap. Sounds varied in pitch (200–1400 Hz; IRN), location (-120–120\degree; HRTF), and pitch/location salience.

Results

Initial fMRI data analysis showed that the auditory tasks (discrimination or memory) modulated activations in supratemporal areas extending from the anterior insula to IPL. As in our previous studies, direct comparisons of the two types of auditory tasks revealed stronger activations in temporal and insular areas during discrimination than memory tasks, while areas in the IPL were more activated during memory than discrimination tasks. Further, pitch tasks were associated with stronger activations than location tasks in the lateral STG/STS, while location tasks enhanced activations in areas of the IPL. During visual tasks, pitch-varying sounds enhanced activations in the lateral STG and HG, while location-varying sounds enhanced activations in more medial areas in the PT.

Conclusion

Activation differences between discrimination and memory tasks were strong and widespread, whereas differences between pitch and location tasks and between visual task blocks with pitch-varying or location-varying sounds were less pronounced. These results demonstrate that activations to spatial and nonspatial sounds strongly depend on the behavioral task and that these activations cannot be explained only by stimulus-dependent processing of physical features of sounds.
Beyond the Auditory Scene: Non-Auditory Predator Cues Modulate Auditory Responses of Single Neurons in the Mouse Amygdala

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Background
Female mice emit a low frequency harmonic (LFH) call in two different contexts, during mating and in response to physical threat or pain. Male mice respond differently to the LFH call in the presence of predator odor or female urine; it is aversive in the presence of cat fur but not in the presence of female urine. The basolateral amygdala (BLA) has auditory inputs from both auditory thalamus and cortex and has been shown to code the valence of social vocalizations. We tested whether there were neural correlates of the behavioral valence change in the responses of BLA units.

Methods
Local field potentials (LFPs) and spiking activity were recorded simultaneously from the BLA of 10 awake, freely moving male mice in response to the LFH call and broadband noise (BBN). Signals were transmitted wirelessly from custom multi-electrode implants using a 16-channel wireless headstage (TBSI Technologies). Responses to sounds were recorded from animals in 4 contexts: clean chamber, neutral condition (cotton wool stimulus), cat fur, and female mouse urine. Responsiveness, for LFPs and spikes, was assessed using t-tests comparing background activity to the post-stimulus activity. A rate modulation index (RMI) was computed during each context to quantify the spike discharge of responsive neurons.

Results
LFPs showed significant auditory-evoked elements for both BBN and the LFH. Responses were typically significantly modulated by both cat fur and female urine and often by the cotton wool bud. Thirty of 64 single neurons responded to the sounds; both excitatory and inhibitory responses were common. The RMI of responses to BBN and LFH changed with acoustic context. The mean absolute RMI value did not change in the presence of cat fur, suggesting that the sounds were equally detectable in each context. However, for individual neurons, the spike discharge patterns changed. The RMI in response to the LFH call increased in the presence of cat fur (Wilcoxon rank, p=0.004); 14 of 30 neurons switched from inhibition or no response to excitation, while 3 of 30 neurons increased their excitation. The RMI of BBN responses decreased in the presence of cat fur (Wilcoxon rank, p=0.025), representing a generalized shift to suppression (15 of 30 neurons) or to greater suppression (5 of 30 neurons).

Conclusion
Auditory responses in the mouse BLA, to both BBN and the LFH call, are differently modulated by non-auditory contextual cues. Supported by NIDCD grant R01 DC 00937.
Retention of auditory memory during breaks in perceptual training.

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Background
What circumstances are required for auditory perceptual learning to occur? One circumstance, indicated by recent findings, is that learning across days requires a sufficient amount of training per day. This suggests information is accumulating over the course of practice and must reach some learning threshold to enable across-day improvement. Interestingly, the memory of this accumulated information appears to decay, because a break in the midst of otherwise sufficient practice can disrupt learning. Here we asked whether this memory begins to decay immediately after practice ceases, as for declarative memory, or instead is retained for some time before decaying. We did so by comparing the ability of two mathematical models, with either immediate or delayed decay, to predict previously reported human learning data.

Methods
Four training groups practiced frequency discrimination (standard: two 15ms 1kHz tones separated by 100ms) for 6-8 days. Each day, groups practiced 360 trials (short-training), 900 trials (long-training), 720 trials with a 30-minute break after 360 trials (30’ break), or 720 trials with a 6-minute break after every 120 trials (5x6’ breaks). A control group only participated in pre- and posttests.

We tested immediate- and delayed-decay models, each with 64,000 parameter settings, by Monte Carlo simulation. For each model and parameter setting, we determined how well the amount of decay corresponded with the amount each group learned.

Results
Only the long-training (p < 0.001) and 5x6’ breaks (p = 0.02) groups showed evidence of learning; the other groups did not (short-training, p = 0.296 and 30’ break, p= 0.423; ANCOVA vs. controls with pretest and day as covariates).

With their best fitting parameters, both models predicted which groups learned and which did not. However, the average performance across all parameters was better for the delayed-than the immediate-decay model (Bayes Factor 7.23). The estimated delay of decay was ~5 minutes.

Conclusion
Auditory perceptual learning appears to require sufficient training per day for learning to occur across days. Our model evaluations suggest that information accumulated during the course of this training is retained in memory for ~5 minutes in the absence of practice. This memory appears to be distinct from declarative memory where decay seems to begin immediately, and from echoic or working memory, which last for less time. The limitations of this memory store place constraints on perceptual training schemes.

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Attentional Mechanisms for Recognizing Acoustic Scene

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Background
Humans exhibit a remarkable ability at processing complex acoustic information from their surroundings and identifying the nature of the environment where they are. This ability is further heightened when guided to detect particular target sounds, even in presence of distracting sounds or ambient noises.

Methods
In the present study, we explore the neural underpinnings of auditory scene recognition in a biomimetic computational model. We exploit the detailed multi-resolution spectro-temporal analysis performed by the mammalian auditory cortex to perform an intricate mapping of sound events present in different acoustic scenes. This framework is further extended by exploring the role of task-driven attention in modulating sensory processing of acoustic information; and ultimately enhancing the ability to detect target sounds of interest.

Results
We demonstrate the ability of the system to outperform previous systems at the task of acoustic scene recognition. We further show the adaptability of the system to recognize particular targets in a noisy environment.

Conclusion
We discuss the relevance of our results in tasks of auditory scene identification and target sound recognition; as well as the role of cortical processes and top-down attentional modulation in sound processing and perception of complex acoustic scenes.

Cortical Representations of Music in Human Listeners

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Background
Cortical neural representations of music in humans are investigated using analysis techniques already demonstrated for analogous cortical representations of speech (Ding and Simon, PNAS 109(29), 11854-11859).

Methods
Using magnetoencephalography (MEG), we present and analyze neural responses to spectrally and rhythmically diverse samples of music. The neural responses are characterized by a temporal response function (TRF) generated via reverse-correlation with the stimuli.

Results
Correlations between actual and predicted responses to musical stimuli are comparable to those obtained using speech stimuli, but unlike the speech case, there is substantial diversity in TRF temporal profiles. The TRFs fall into at least two categories based on the polarity of the earliest peak. Critically, pairwise correlations of TRFs between subjects for each stimulus are generally higher than pairwise correlations of TRFs between stimuli, indicating that the encoding mechanism is robust across subjects.

Conclusion
This suggests that the underlying mechanism for encoding music in the brain is feature-dependent, and may specifically depend on the rhythmicity of the music.
A Neuromechanistic Model of Auditory Streaming
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Background
Interleaved tone sequences (AB_AB,... and ABA_ABA,...) are commonly used to study auditory streaming. For ranges of A-tone and B-tone frequency differences (DF) and presentation rate (PR) stimuli are ambiguous, leading to different percepts, integration or segregation with bistability and alternations between integration and segregation during long presentations. Neurophysiological results for A1 support a two-dimensional feature mapping (tonotopy and periodotopy) and reveal neuronal dynamics consistent with integration-segregation switching. While phenomenological models for perceptual bistability exist, the development of neuronally-based models is immature.

Methods
We have developed idealized firing-rate models with a 3-population recurrent network architecture for the ‘percept formation’ stage (see Figure). Inputs to the network embody the tonotopic and periodotopic representation in A1, e.g., population A responds most to the A-tone while population AB is most responsive to frequencies midway between A-tone and B-tone frequencies. If A and B are alternately activated by the interleaved sequence with little activity in AB (say for large DF) we say that segregation is being perceived. Integration occurs when AB is dominant (or equi-active with A and B), say for small DF. Activation is partially retained in the silent gaps between tones by recurrent excitation with slow, NMDA-like, facilitatory synapses. Competition for percept dominance occurs by lateral inhibition and slow negative feedback from firing-rate adaptation underlies integration-segregation alternations.

Results
First, we show that by implementing tonotopy and periodotopy in an A1 model we can account for: 1) experimentally observed A1 neuronal responses to interleaved A and B tones and 2) differential suppression phenomenon i.e., frequency selectivity of A1 neurons improves with repetition of AB sequences (Fishman et al, 2001). For a range of input and coupling parameter values, the model shows coexistence between the integration and segregation states – bistability, consistent with stimulus ambiguity. When adaptation is strong enough, the model produces regular transitions between these two coexisting states. The percent of time that integration or segregation is dominant depends on the input values.

Conclusion
Our model provides a neuronally-based rationale for specifying inputs to a percept formation network, enabling us to model the effects of changing DF and PR. While bistability has been modeled for the visual case (say, binocular rivalry) inputs are typically constant in time. The dynamical aspects of the interleaved auditory stimuli (with silent gaps between tones) present new challenges for modeling bistable perception that we have overcome by utilizing synaptic dynamics (NMDA-like slow decay and facilitation) to avoid re-setting activity variables between tones.
Receptive field changes in Primary and Secondary Auditory Cortex as correlates of streaming in behaving ferrets

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Background
Alternating two-tone sequences are usually perceived as two streams when the tones are far apart, while synchronous two-tone sequences evoke a unified single stream percept regardless of their separation. Consequently, it was hypothesized that the receptive fields (RFs) in auditory cortex would be different when listening to these two types of sequences, with the latter inducing broadly tuned receptive fields compared to the former.

Methods
Ferrets were trained to listen attentively to the two-tone sequences until they transitioned to a random sequence of tones, at which point they were to approach a spout for a water reward. Responses were recorded in the primary auditory cortex (A1), and also in an adjacent secondary auditory field on the posterior ectosylvian gyrus (PEG) during behavior and in the passive listening. For the tone sequences, one tone (B-stream) was usually fixed at a neuron's BF, while the other tone (A-stream) varied between 0.25 to 2 octaves away from the first tone in different trials. The final random sequence allowed us to measure the neuron's RFs at the end of the two-tone sequences.

Results
In A1, both two-tone sequences evoked phase-locked responses that rapidly adapted after the onset of the trial, reaching a steady state level after 3-4 tones. When the animal was attentively listening, however, the responses to the two types of sequences diverged considerably. The alternating tone responses became more suppressed relative to the passive listening. By contrast, responses to the synchronous sequence were enhanced. This difference in responses was also evident in the RFs of the neurons at the end of each sequence. Since the alternating and synchronous trials were randomly presented, the results suggest that these RFs differences built up rapidly within each trial that typically lasted between 1-3 seconds. Preliminary responses from the PEG reveal a somewhat different picture, with weaker phase-locking and far noisier RFs, making similar response measurements to the A1 very difficult.

Conclusion
In summary, we believe that response changes such as those in A1 are mediated by rapid plasticity in the connectivity among the responding neurons. Specifically, we hypothesize that during behavior, stimulating neurons incoherently induces mutually suppressive connectivity, while the opposite is true for coherent stimulation. These patterns of RFs change are consistent with the idea that stimulus coherence is the key in inducing the streaming percepts and the organization of complex auditory scenes.
Task-dependent spatial responses in auditory cortex of common marmosets
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Background
Auditory cortex is essential for behaviors involving sound localization in mammals. Although spatial processing in auditory cortex has been extensively studied in passive subjects, there are few studies of spatial processing under behaving conditions. In this study, we investigated how representation of space by auditory cortex neurons is modulated as specific regions of space become behaviorally relevant.

Methods
We recorded single-unit responses in auditory cortex of marmosets (Callithrix Jacchus), an arboreal New World monkey, while they performed a spatial discrimination task with either a contralateral or ipsilateral stimulus bias. We asked whether any observed changes in response properties would depend on either the spatial parameters of the task, or the spatial tuning properties of the neurons themselves.

Results
We found a population of neurons for which firing rates to target locations recorded during task engagement were increased when compared to those recorded during passive listening. There was also a smaller increase in firing rates to target locations in hit trials compared to miss trials. A subset of neurons showing increased firing rates to targets also had increased firing rates to background (non-target) locations. Neurons which showed increased firing in response to one target location tended to have increased firing rates to other target locations as well, and neurons with increased firing rates in the one behavior condition (e.g. contralateral stimulus bias) tended to also have increased firing rates in the opposite condition. Furthermore, neurons with increased firing rates included those preferring both contralateral and ipsilateral locations in the passive condition.

Conclusion
Taken together, these results showed that responses of a subset of neurons in auditory cortex are modulated by spatial tasks.

Laminar Recordings of Task-Related Receptive Field Plasticity with Multisite Depth Electrodes in the Primary Auditory Cortex (A1) of the Behaving Ferret
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Background
Previous studies have shown task-related receptive field plasticity (RFP) in A1 neurons (Fritz et al., 2003) during performance of simple spectral discrimination tasks. Two open questions were: (1) whether adaptive RFP changes also occurred during performance of temporal discrimination tasks, (2) whether such RFPs were equiprobable for A1 neurons in all cortical layers, or located preferentially in certain layers or cell types.

Methods
We trained ten ferrets to perform two auditory discrimination tasks in which they learned to discriminate between broadband rippled noise stimuli and narrow-band tones (spectral task), and/or between low and high-rate click trains (temporal task). The basic behavioral paradigm for both tasks was conditioned avoidance (Fritz et al., 2003). We recorded from behaving head-fixed ferrets, using simultaneous multichannel recordings, and demonstrated plasticity at a single-unit and population level for temporal tasks. To explore the question of RFP laminar specificity, we recorded from A1 in three trained ferrets during behavior using near-perpendicular penetrations with multisite depth electrodes that allowed for simultaneous recording from all cortical layers, and comparison of RFP in each layer. We reconstructed the laminar location by using CSD (current source density) analysis on the LFP from all sites. We compared the STRFs (spectrotemporal receptive fields) recorded during a pre-behavior quiescent state and then during active discrimination behavior in all neurons.

Results
Many neurons showed STRF plasticity along the time axis during performance of temporal tasks. To help clarify whether these changes were specific for temporal tasks, or whether they might also occur in spectral tasks, we recorded from A1
neurons under both task conditions. We found STRFs throughout the depth of the electrode recording sites, and ~65% of all STRFs showed task-related plasticity for either spectral and/or temporal tasks. Our preliminary data suggests greater STRF plasticity occurs in supragranular layers. Some plastic cells showed RFP for only one task condition. About 60% of plastic neurons showed RFP for both tasks, however, the effects were often different. We also observed task-related changes in neuronal responses to task stimuli that were not captured by the STRF which were found in all layers.

Conclusion
A1 neurons demonstrate remarkable, task-related RFP for both spectral and temporal tasks. Our results give first insights into the laminar location of this type of attention-driven RFP, and also indicate that the STRF adapts to task conditions along both the time and frequency axis.

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Comparison of Neuronal Responses in Ferret Frontal Cortex during Positive and Negative Versions of an Auditory Long-Term Pitch Memory Task.
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Background
An earlier study on ferret frontal cortex (FC) revealed persistent neuronal responses to auditory target stimuli over a time course of minutes to hours (Fritz et al., 2010) following performance of auditory discrimination tasks. In order to explore the role of the FC in auditory long-term memory, we developed a new auditory task that requires information retrieval from long-term pitch memory.

Methods
Six ferrets were trained to classify single tonebursts into three frequency ranges (Low, Middle, High). Three animals were trained on a positive reinforcement (PR) task version, in which they learned to lick a waterspout for reward (Go-response) when the toneburst fell in the middle frequency range, and to stop licking the waterspout after tonebursts in either the low or high frequency range (NoGo response). In the PR task, licking during Nogo stimuli lead to a timeout. Three additional ferrets were trained on a conditioned avoidance (CA) version of the task, in which they received mild shock if they licked to tones in the middle range (Nogo stimuli), but could lick freely through the tones in both low and high range (Go stimuli). Each trial simply consisted of the presentation of a single toneburst, and the animal’s response. Each daily session consisted of 100-200 trials. After animals were trained, they were implanted with headposts to secure head position during task performance. We recorded chronically from the FC of behaving, headfixed ferrets, using an independently-moveable multiple electrode system.

Results
All ferrets learned the basic discrimination, with either training paradigm (PR or CA), in an average of one month. Moreover, although toneburst stimuli in the initial training set had wide frequency spacing, animals generalized and readily transferred to a denser stimulus set with new frequencies, after learning the original task. Preliminary neurophysiological results from the PR task indicate the presence of two neuronal classes in FC that were strongly activated during task performance and selectively preferred tonal frequencies that corresponded to either \(\frac{1}{2}\)Go\(\frac{1}{2}\) stimuli or \(\frac{1}{2}\)Nogo\(\frac{1}{2}\) toneburst stimuli (Yin et al 2012). We will compare neuronal response from FC during the CA version of the task to those obtained in FC of animals trained on the PR version of the task.

Conclusion
These preliminary results provide insights into how the FC encodes, represents, classifies and retrieves the associative meaning of sensory stimuli during performance of an auditory long-term memory pitch task under different valence conditions.

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Human Cortical Auditory Evoked Offset and Onset Response Sensitivity to Signals in Noise
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Background
Cortical auditory evoked potentials (CAEPs) can be elicited by both the onset and the offset of an auditory stimulus. However, in both the clinical and research literature, offset responses have been both underreported and understudied in
comparison to onset responses. Furthermore, while single unit animal studies have demonstrated important differences in the specific characteristics of the neural populations responsible for the onset and offset response, far-field studies (EEG and MEG) have tended to suggest that onset and offset responses are not physiologically independent.

The gap in resolution between single unit and far-field recordings is significant enough that, despite certain observable differences in the characteristics of the neural populations that respond to onsets and offsets, the usefulness of offset CAEPs in the neurophysiologic characterization of auditory processing and processing disorders remains to be convincingly demonstrated. The goal of studying offset responses in addition to onset responses then, is to determine how and when offset responses differ from onset responses such that additional information about the encoding of the stimulus may be obtained.

We report that offset responses are sensitive to changes in absolute signal level while SNR (signal-to-noise ratio) is held constant.

**Methods**

Billings et al. (2007): Responses were recorded from 13 young normal-hearing subjects, who were presented a 1-kHz tone (757 ms) at 7 signal levels (range: 30 to 90 dB SPL).

Billings et al. (2009): Responses were recorded from 15 young normal-hearing subjects, who were presented a 1-kHz tone (757 ms) at 2 signal levels (60 and 75 dB SPL).

**Results**

We find that, while peak latency and amplitude of onset responses are not sensitive to changes in absolute signal level if signal-to-noise ratio is held constant, offset latencies and amplitudes are, while also being sensitive to changes in SNR.

**Conclusion**

Cortical Auditory Evoked Offset responses are not simply less salient versions of onset responses, but can be measurably different even in far field recordings. In the present findings, offset responses indicate that the cortex is sensitive to absolute signal level cues in the encoding of signals in noise, a sensitivity not apparent in onset responses.

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**Bold Response in the Auditory Cortex of Non-human Primates Matches Pitch Percept across Different Repetition Rates**

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¹Newcastle University

**Background**

Pitch is a fundamental percept with a complex relationship to basic acoustic stimulus properties such as frequency composition and temporal regularity of sounds. Thus a neuronal correlate of pitch is best identified by looking for neural responses that match the varying pitch percept to changing stimulus attributes.

**Methods**

Here we measured the BOLD response across the lower and upper limit of the pitch percept to regular interval noise (RIN) in the auditory brain upstream from the inferior colliculus (IC) of two rhesus macaques. The data were recorded in a Bruker 4.7 T vertical scanner during fixation of a visual stimulus. In three different experiments we presented RIN stimuli at repetition rates from 8 Hz to 2048 Hz and we used fixed-amplitude random-phase broadband noise as control.

**Results**

In both animals, widespread areas in the auditory cortex, particularly in lateral and anterior core and belt, showed a progressive increase in the BOLD response to increasing repetition rates beginning at the lower limit of pitch in humans of about 30 Hz. This effect was most pronounced bilaterally in a region between the core areas A1 and R and the adjacent belt areas. The response increase levelled off in a plateau between 256-512 Hz. Responses in the IC and the medial geniculate body (MGB), while showing stronger response to repetitive sounds than noise, did not show a modulation of the response across the lower pitch level. At 1024 Hz and above, coinciding with the upper limit of pitch to RIN in humans, all areas in the auditory cortex, MGB and IC showed a progressive decay in the BOLD response.

**Conclusion**

Macaques show a response profile in the auditory cortex that matches the pitch percept across the lower and upper human pitch limit. Behavioural tests in macaques suggest a similar lower pitch limit than humans [1]. The effect is
consistently strongest in an area that has previously been highlighted as a “pitch center” in marmosets [2], supporting its role a candidate for a neural correlate of pitch. These findings are in line with results from single unit recordings in macaques [3]. Responses in the IC and the MGB across the lower pitch limit do not match perception, suggesting that the pitch percept arises at the level of the cortex.


A modeled auditory evoked potential predicts attentional modulation of averaged and single trial EEG signals
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¹Boston University

Background
Human listeners easily can focus attention on a target sound within a complex auditory scene, though the underlying mechanisms of auditory attention are still largely unknown. Attention is known to modulate the cortical representation of the auditory scene, an effect that alters auditory evoked potentials (AEPs). These AEPs consist of different components with different latencies originating from different brain locations. Observing how attention modulates different AEP components can help us understand which neuronal centers have roles in selective attention.

Methods
Three competing isochronous tone sequences were presented (left, center, and right). Each stream had a different rhythm. AEPs were measured while listeners selectively directed attention to either the left or right competing stream. The AEPs from different attentional conditions were compared. A model based on a superposition of Gaussian mixture functions convolved with a train of pulses with a pulse at each stimulus onset was used to generate predicted AEPs. By finding the model parameters that achieve the best match with measured AEPs, we derive a quantitative measure of attentional modulation.

Further, we analyzed single trial EEG results to predict which stream was attended on that trial, based on template-matching pattern classification. Different AEP components were incorporated into the templates, in order to compare which components were most useful in determining which stream a listener attended.

Results
Measured AEPs showed strong attentional modulation. The best fitting models attenuate AEP components from the ignored streams by 20dB, while evoking strong P1, N1, and P2 components at onset events in the attended stream. The highest correlation between the measured AEPs and modeled was 0.80, averaged over mid-frontal 22 channels. P1 and P2 components were important for predicting the averaged AEP shapes, but had no influence on single-trial classification of which direction a listener was attending. In contrast, the model containing only the N1 component (“Model C”) achieved the best classification accuracy: 72%, for a 3-second-long single trial.

Conclusion
Attention strongly modulates the cortical representation of a complex auditory scene. This modulation causes the AEPs in response to an ignored stream to be attenuated by 20dB, on average. The N1 component is the strongest marker for predicting the direction of a listener’s attention from a single trial. The proposed modeling framework can be a useful tool for investigating mechanisms underlying auditory attention, and can provide a feature basis for a brain-computer interface paradigm using EEG and auditory stimuli.
Late Auditory Evoked Potentials in Responses to Informational Masking Stimuli
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1Hearing and Speech Laboratory, University of California Irvine, 2Center for Hearing Research, 3Dept. of Otolaryngology, University of California Irvine

Background
The auditory system creates a neuronal representation of the acoustic world based on cues present at the listener’s ears. Parsing individual sources becomes challenging when multiple sources are active. The current study aims to understand the neuronal mechanisms underlying this process known as auditory scene analysis (ASA). Here, we measured late auditory evoked potentials (LAEPs) in response to complex signal-in-masker stimuli designed to elicit informational masking (IM). Published psychophysical data show large inter-subject variability in response to these IM stimuli. Our aim is to develop a LAEP-based paradigm that could test whether this large variability is detectable in neuronal responses.

Methods
Stimuli consisted of a five-component tone complex, 50 ms in duration, repeated four times at a rate of 10 Hz (total duration 350 ms). The signal was one component fixed at 4000 Hz, and the four masker components either varied in frequency across bursts or were fixed in frequency. Signal and masker were gated synchronously. All energetic masker (EM) frequencies were drawn from a ±1/3-octave band centered at signal frequency, whereas IM frequencies occupied a broad range outside of the signal band. We obtained mismatch-negativities (MMNs) using an optimized oddball paradigm: One standard (masker alone), randomly interleaved with five deviants (signal + masker). Deviant signal levels were 30, 50, 60, 70, and 90 dB SPL. Masker components were fixed at 60 dB SPL. Subsequently, subjects performed a psychophysical 2-AFC detection task with an adaptive tracking procedure.

Results
In IM conditions, MMN components of all three subjects decreased systematically with decreasing signal level. In contrast, for EM only the highest signal level elicited significant MMNs. Accordingly, MMN thresholds were lower for IM (30–50 dB SPL) but not for EM (70–90 dB SPL). In accordance with previously reported psychophysical data, neural thresholds displayed large inter-subject variability for IM. In one subject we tested psychophysical thresholds and found a strong positive correlation with neuronal thresholds (R2 = 0.99 N = 4 p < 0.001).

Conclusion
We demonstrated that 1) it is possible to record meaningful LAEPs for these relatively long and complex stimuli 2) MMNs systematically vary with subjective task difficulty and 3) where available, neuronal thresholds correlate well with behavior. Although more data are needed to solidify these findings, our experiments potentially provide an important insight into the relationship between behavior and cortical activity during ASA.
Neural Correlates of Musical Discourse: an fMRI Study of Interactive Jazz Improvisation
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¹University of Pennsylvania, ²Johns Hopkins University School of Medicine

Background
Interactive generative musical performance provides a suitable model for communication because, like natural linguistic discourse, it involves an exchange of ideas that is unpredictable, collaborative and emergent.

Methods
We utilized functional magnetic resonance imaging to investigate a form of jazz improvisation called trading fours, during which two musicians participate in a call and answer exchange akin to a ‘musical conversation’. Eleven professional jazz piano players participated in this study. Two experimental paradigms were used during scanning, one of low complexity and one of high complexity. In the low complexity task, the subject and experimenter either traded a one octave, ascending and descending D Dorian scale (control task) or traded four measure improvisations (experimental task). Improvisation was heavily restricted to continuous, monophonic quarter notes in the key of D Dorian. In the high complexity task, the subject and experimenter either traded four measures of a memorized jazz composition (control task), or traded four measure improvisations (experimental task). All tasks were performed with accompaniment from a prerecorded rhythm section.

Results
SPM8 was used to analyze all functional neuroimaging data. Imaging data contrast analyses of improvised trading conditions vs. memorized trading conditions revealed activation of perisylvian language areas linked to processing of syntactic elements in music, including inferior frontal gyrus and posterior superior temporal gyrus. Furthermore, this form of improvised musical communication was associated with significant deactivation of angular gyrus and supramarginal gyrus, brain structures directly involved with semantic processing for language.

Conclusion
Perisylvian language areas that subserve syntactic function are active during improvised musical communication. These findings support the idea that the exchange of ideas in music does not involve traditional semantic processing. Furthermore, these findings suggest that neural regions for syntactic processing may be domain-general for communication rather than domain-specific for language.
### Activations

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Juvenile Auditory Learning is Associated with Diminished Cortical Synaptic Inhibition

Emma Sarro¹, Vibhakar Kotak¹, Dan Sanes¹
¹New York University

Background
Auditory associative training that occurs during development leads to improved performance in adulthood (Sarro and Sanes, 2011), indicating that developing animals display a unique benefit from sensory experience. To test whether functional adjustments in primary auditory cortex accompany such learning, we assessed inhibitory synaptic function, which has been implicated in cortical plasticity (Froemke et al., 2007).

Methods
Juvenile gerbils were trained on an amplitude modulation detection task from postnatal day (P) 23 to 40. Subsequently, we generated thalamocortical brain slice preparations on the final day each animal received training, and assessed inhibitory synaptic function. Spontaneous inhibitory synaptic currents (sIPSCs) were recorded in layer 2/3 pyramidal neurons of thalamorecipient auditory cortex following blockade of ionotropic glutamate receptors and sodium spikes.

Results
Auditory associative learning was accompanied by an immediate reduction in both the amplitude and frequency of sIPSCs, and this decrease persisted for several days. However, following longer training durations, sIPSC amplitude and frequency approached or exceeded control levels. Finally, those juveniles that did not display significant learning, also showed control-like inhibitory currents.

Conclusion
Thus, a limited period of diminished inhibition may facilitate associative learning in developing animals.
The Effects of Reward Expectation on Neural Responses in Primary Auditory Cortex (A1) During the Learning and Performance of a Tone Detection Task
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¹University of Maryland

Background
The neural network underlying reward and reward expectation has not traditionally included primary sensory cortices. However, recent studies in behaving animals have shown that neurons in A1 can encode target expectation and that neurons in V1 (primary visual cortex) can encode reward timing. To explore the effects of reward expectation on A1 coding of target sounds, we studied amplitude-modulation (AM) coding of target tones in A1 during task conditions involving expectation of a water reward.

Methods
Two ferrets were trained on a conditioned avoidance (CA) tone detection task. In each trial, a series of rippled noise stimuli was followed by a 4 Hz AM tone. During each testing session, a different AM carrier (ie. target) frequency was used. Animals were trained to lick from a waterspout for 1.5 seconds after AM tone offset, but to refrain from licking during a variable duration (1-4 second) target AM tone to avoid a mild tail-shock. Blocks of trials in a given session had fixed duration tones allowing the animal to accurately anticipate reward timing. While a ferret was learning the task, we recorded single-units in A1 using a moveable 64-channel multi-electrode array in a behaving, head-fixed ferret to study the plasticity of A1 responses to the task sounds over the time course of learning (~1 month).

Results
During task performance in the trained animal, the spike-rate steadily increased over the duration of the target AM tone. The synchrony of spiking to the 4 Hz AM was decreased during the task. Consistent with previous findings (Fritz et al, 2003; Atiani et al., 2009), we also saw an overall suppression of spike-rate during the CA task, and persistent task-dependent frequency-specific changes in the shape and gain of spectro-temporal response fields (STRFs).

Conclusion
Our results suggest that reward anticipation in combination with inhibited reward-seeking behavior in the CA task leads to a build-up of A1 responses to target sounds as the reward time approaches. The decreased AM spike synchrony during the task reflects, in part, a combination of the decreased AM-feature salience for successful task performance, the suppressed overall spike-rate during the task, and an increase in the jitter of spike timing across trials during behavior. We will further discuss task-dependent effects on STRF tuning, and the effects of learning the task on A1 responses to the task sounds during learning.
Dynamics of plasticity in human auditory cortex during course of perceptual learning
Serin Atiani¹,², Robert J. Zatorre¹,², Marc Schönwiesner¹,³
¹Montreal Neurological Institute, ²McGill University, ³Université de Montréal

Background
Establishing the link between behavior and its underlying neural mechanism is one of the central goals of neuroscience. Probing the representation of behaviorally significant sensory experiences is critical to understanding the functional dynamics of our sensory systems. The significance of the auditory system coupled with humans’ unique ability to learn and adapt makes the human auditory system well suited to studying these questions. In this study we manipulate the behavioral significance of stimuli through training participants on an acoustic task and observe the changes in their representation that take place over the course of learning. Our experiment is designed to take advantage of auditory neurons’ selectivity to spectro-temporal features of sound (Klien et al 2000, Theunissen et al 2000, Depireux 2001, Schönwiesner&Zatorre 2009). We use dynamic ripples, complex broadband stimuli with a drifting sinusoidal spectral envelope (Kowalski et al 1996), to measure modulation transfer functions (MTF) for voxels in auditory cortex, while participants are actively listening to a series of these ripples as they perform the task.

Methods
Twelve participants were presented with a series of dynamic ripple stimuli and were trained to identify a target ripple with a unique spectro-temporal combination from a set of 16 different ripples with 4 spectral and 4 temporal modulation rates. Training took place during 5 consecutive days. Functional brain images were acquired with a 3T MRI scanner on the first, second and fifth day during task performance. We used the ripples heard during the task to map spectro-temporal (MTF) of voxels in auditory cortex (method described in: Schönwiesner&Zatorre 2009).

Results
Participants’ performance improved through the course of training to reach optimal performance by the fifth day. We observe task related changes in the MTF of auditory cortex voxels starting on the first day of training reflecting a clear selectivity for the target ripple. This selectivity improves during the course of learning, by continued differential response to the target ripple coupled with gradual drop in response to remaining ripple set, that is accompanied by a drop of global BOLD signal in auditory cortex.

Conclusion
Our study demonstrates changes in spectrotemporal modulation tuning during learning that reflect the modulation content of the target sound. Preliminary analyses suggest a correspondence between cortical changes in tuning and behavioral changes in target.
Understanding the Effects of Central and Peripheral Masking in Cochlear Implant Users
Carol Q. Pham1,2, Peter Bremen3, John C. Middlebrooks2,3, Fan-Gang Zeng1,2, Myles Mc Laughlin1,2
1Hearing and Speech Laboratory, University of California, Irvine, 2Center for Hearing Research, University of California, Irvine, 3Department of Otolaryngology-Head and Neck Surgery, University of California, Irvine

Background
Although cochlear implant (CI) users communicate relatively well in quiet, they have great difficulty understanding speech-in-noise. Previous studies in normal hearing (NH) subjects have demonstrated that both peripheral and central processes play a significant role in speech-in-noise perception. While it is known that wide peripheral filters in CI users contribute to poor speech-in-noise performance, the role of altered or degraded central processing has been less well studied. Here, by first measuring peripheral filters, we could use carefully controlled informational (IM) and energetic masking (EM) stimuli to study the roles of central versus peripheral processes, respectively, in electric hearing.

Methods
Four post-lingually deaf Nucleus 24 users were tested. Peripheral filter bandwidth was measured using standard electrically evoked compound action potential forward masking techniques. All stimuli were presented through a research interface (HEINRI) at a rate of 300 pulses per second. The signal was fixed on electrode 11. The masker was on four other electrodes placed either inside (EM) or outside (IM) of the peripheral filter. Both signal and masker were 40 ms bursts, pulsed 4 times at 5 Hz. Maskers with synchronous and asynchronous onset time re signal were tested as were maskers with varying (V) and constant (C) electrode position over time. Masker level was roved to avoid introducing any loudness cues. A two-interval, forced-choice adaptive procedure was used to measure signal detection thresholds in the presence of maskers.

Results
Surprisingly, the amount of masking produced by stimulating electrodes outside of the peripheral filter (IM) tended to be greater than that produced by stimulating electrodes inside of the filter (EM). In agreement with previously reported data in NH listeners, CI data showed that (1) thresholds with synchronous maskers were higher than those obtained with asynchronous maskers, and (2) thresholds for C maskers were higher than thresholds for V maskers. In general, inter- and intra-subject variability was large for all masker types.

Conclusion
Results indicate that in electric hearing the effects of central masking can often be greater than peripheral masking. While spectral cues did improve signal detection for some CI subjects, the effect of temporal cues dominated and improved signal detection for all subjects tested.

A Spectrally Rippled Noise Mismatch Negativity Paradigm for Objectively Assessing Speech Perception in Cochlear Implant Users
Myles Mc Laughlin1,2, Alejandro Lopez Valdes2, Laura Viani3, Peter Walsh3, Jacklyn Smith3, Richard B. Reilly2, Fan-Gang Zeng1
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Background
A number of recent studies suggest that cortical evoked potentials (CEPs) may provide a useful objective measure of speech perception in cochlear implant (CI) users. However, weak correlations with behavior and artifact cancellation issues have hampered the clinical impact of CEP based objective measures in CI users. Psychoacoustic studies have shown that a CI user’s ability to discriminate spectrally rippled noise stimuli correlates reasonably well with their speech perception. Here, we have developed a clinically applicable single-channel artifact cancellation approach which allows the acquisition of artifact free CEPs using complex stimuli. Using this approach we demonstrate how CEPs to spectrally rippled noise may be used to estimate speech perception in CI users.

Methods
We developed a high bandwidth (0-100k Hz), high sample rate (125kHz) CEP acquisition system which allows us to clearly resolve the CI related artifact. When acquired with this system the artifact is completely high frequency (>= CI stimulation rate) and can be removed using standard filtering techniques. Using this system we measured mismatch negativity (MMN) responses in 10 CI subjects and 10 normal hearing controls. The standard stimulus (90%) was spectrally rippled broadband noise; the deviant (10%) was the inverted version, having an equal number of ripples per octave (RPO). To define a neural RPO detection threshold we increased the number of RPOs until the MMN response
was no longer greater than a noise floor determined from the standard only responses using a boot-strap method. Behavioral RPO detection thresholds were also measured using a two alternative forced choice paradigm.

Results
The high bandwidth, high sample rate system provides robust, fully automatic, real-time, CI artifact removal. A significant correlation (p=0.02, r²=0.54) was found between the neural RPO detection threshold and the behavioral RPO detection threshold.

Conclusion
Using a single-channel clinically applicable artifact removal technique it was shown that CEPs can be used to estimate the spectral processing abilities of CI users. Given the known correlation between speech perception and the behavioral RPO detection threshold, we expect to see a correlation with neural RPO detection threshold and this is currently being tested.

Temporal Processing Properties of the Auditory Evoked Potentials in Cochlear Implant Users
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Background
For perception purposes, the auditory system encodes temporal and spectral information of sounds. The primary purpose of this project is to investigate the temporal properties of auditory evoked potentials (AEPs) at the peripheral and cortical stage of the auditory system in cochlear implant (CI) users. This research may lead to future efforts toward improving the temporal representation of sounds, thereby maximizing CI performance. We expected the temporal properties of the late auditory evoked potential (LAEP) from the cortex to be related to that of the electrical compound action potential (ECAP) from the auditory nerve, given that the temporal properties may be kept and/or accumulated after multiple neural synapses along the auditory system.

Methods
A group of postlingual CI adults were recruited. A stimulus-train paradigm was used to examine how the AEPs change over time. The ECAP was recorded using NRT software (Custom Sound EP 2.0) installed on the clinical fitting system. Since the NRT software does not allow for ECAP recording in response to individual pulses within a pulse train, the approach described previously was used (Hay-McCutcheon et al., 2005). Specifically, the response to a probe that is preceded by a varying number of a series of masker pulses was measured. Offline subtraction was performed to extract the ECAP to each pulse in a pulse train (1000 pps, 100 ms). The LAEP was recorded using the 40-channel Neuroscan system. Tone bursts at 1 kHz were presented according to the stimulus-train paradigm consisting of 30 trains (inter-stimulus-interval = 0.7 ms; inter-train-interval =15 sec) via a loudspeaker placed at ear level, 50 cm from the test ear at 90° azimuth.

Results
Pilot results showed that, for both AEPs, the response was the largest in response to the first stimulus and declined fast until it reached asymptote, displaying adaptation. The adaptation index (AI) was calculated using the formula, AI = Aasymp/A1. The adaptation index ranged from approximately 0.22-0.57 (M=0.42, SD=0.16) for ECAP and 0.19 to 0.72 (M=0.45, SD=0.17) for the N1-P2 complex of the LAEP.

Conclusion
This study is ongoing. Full results will show the correlation between LAEP and ECAP adaptation in CI users and LAEP data from normal hearing controls.

Behavioral Measures of Temporal Processing and Speech Perception in Normal Hearing and Cochlear Implant Listeners
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Background
Although all cochlear implant (CI) users achieve some level of hearing sensitivity, post-implantation speech perception outcomes vary greatly. Previous studies reported a strong correlation between CI users’ phoneme recognition scores and modulation detection thresholds using an electric pulse train directly presented via the electrode array, suggesting that
temporal processing may largely contribute to speech perception (Fu, 2002). The purpose of this study was to examine the relationship between speech perception performance and behavioral measures of temporal processing using the Random Gap Detection Test (RGDT, Keith, 2000) presented in the sound field. The results can help improve understanding the large variability of speech perception performance.

**Methods**
A group of NH subjects (n=10) and postlingually deafened CI users (n=5, 2 bilaterally implanted) participated. Stimuli were presented monaurally via loudspeaker at the most comfortable loudness level (MCL) for CI subjects and insert earphones for NH subjects. The contralateral ear was plugged throughout the entire testing session. The stimuli consisted of the following: Speech Recognition Threshold test, Central Institute for the Deaf (CID) W-22 Word Discrimination test, Banford-Kowel-Banch Speech In Noise (BKB-SIN) test, and RGDT Test.

**Results**
Pilot results showed that the RGDT thresholds are significantly higher in cochlear implant users (26.67 ms, SD = 3.33) than in normal hearing (NH) listeners (7.11 ms, SD = 4.37, p <0.01). The SRT is significantly poorer in CI users (Median=30 dBHL) than in NH listeners (Median=0 dBHL, p <0.01). Discrimination percent correct was significantly poorer in CI users (Median=68%, p <0.01) than in NH listeners (Median=100%, p<0.01). The SNR (signal-noise-ratio) for BKB-SIN test is significantly higher for CI users (M=11.69, SD=4.23) than for NH listeners (M=-4.38, SD=2.70), indicating that the CI subjects required a more favorable SNR in order to be able to repeat 50% of target words correctly. Regression analysis for CI data showed a nearly significant correlation between the mean gap detection threshold and the discrimination score (R2=0.56, p=0.05). There was no significant correlation between the mean gap detection and SRT (R2=0.15, p>.05), or the BKB-SIN results (R2=0.37, p>0.05).

**Conclusion**
This study is ongoing. Pilot results suggest that cochlear implant users do have temporal processing limitation, which is the basis of poorer speech discrimination.
Processing of coherent temporal level fluctuations in Cochlear Implant users

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Background
The normal hearing (NH) auditory system has elaborated strategies to segregate different sounds with overlapping spectra occurring at the same time – a usually unsolvable task for cochlear implant (CI) users. An important neural mechanism in this context is across-frequency processing: the comparison of temporal level fluctuations across auditory filters. Many natural sounds including speech provide such correlated patterns of intensity changes across-frequency that enable comodulation masking release (CMR) and affect auditory stream formation.

Methods
The experiment followed a flanking-band paradigm for CMR measurements using the Nucleus Implant Communicator: uncorrelated and comodulated biphasic pulse trains were addressed to selected intra cochlear electrode contacts (masker components). A steady state level variable pulse train was added to a medial channel (target signal). Masked detection thresholds (MDT) of the target signal were determined according to an adaptive 3-AFC paradigm.

Results
The measured mean CMR was strongly reduced in CI users (3.6 dB ±5.9, p<0.05) compared to NH listeners (12.9 dB ±2.6, p<0.01). However, results of CI users were heterogeneous: 66% of test subjects achieved no (0.1 dB ±2.7) and 33% a high CMR (10.7 dB ±3.2).

Conclusion
We looked for reasons and indicators for CMR in individual CI users. Additional masking as the main reason for the effect is unlikely: the MDT in the comodulated test condition in test subjects with CMR decreased, while MDT in the uncorrelated test condition was not higher than the mean level across all test subjects. These findings point out to a centrally mediated CMR in our study with direct stimulation. Etiology revealed to be a good indicator for CMR: CI users with e.g. acute hearing loss or other sensorineural hearing impairment achieved much higher magnitudes than listeners with e.g. long-term progressive hearing loss or late-implanted congenital hearing loss. Also the duration of hearing loss pre-implantation seemed to be a good indicator for CMR.

The outcome that a subgroup of CI users benefits from amplitude comodulation inspires the development of new signal processing strategies for CI systems in terms of enabling across-frequency processing in everyday settings.

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Recovery from forward masking in cochlear implant listeners depends on stimulation mode
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Background
Bipolar and monopolar stimulation in cochlear implants result in different detection thresholds, suggesting that different groups of neurons may be activated by the two stimulation modes. It is likely that one group (ie that excited by bipolar stimulation) is a subset of the other. Given the recent interest in more focused stimulation of the auditory nerve in cochlear implants, it is important to quantify the response properties of more focused groups of neurons. The objective of this study is to compare the time course of recovery from forward masking in cochlear implant listeners using bipolar and monopolar stimulation modes.

Methods
The masker is a 300-ms long train of biphasic current pulses (100 microseconds/phase, 1000 pulses/s). The probe is a 20-ms long train of identical pulses. Recovery functions are measured under the following three conditions: A. Monopolar masker, monopolar probe; B. Monopolar masker, bipolar probe; C. Bipolar masker, bipolar probe. Stimuli are delivered using a custom research interface and directed to different electrodes across the device. Masker and probe are always directed to the same active electrode, regardless of stimulation mode. The masker is presented at a fixed level, and maskers on different electrodes are loudness-balanced to each other. Psychophysical detection thresholds for the probe were measured using a 2 interval forced choice adaptive procedure. Thresholds were obtained at masker-probe intervals ranging from 2 to 128 ms.

Results
Results suggest that in conditions B and C, recovery functions are nonmonotonic, consistent with previous results obtained with stimuli similar to those in condition C by previous investigators. Condition A seems to result in smoother and faster rates of recovery. Recovery functions for condition A appear to be fairly homogeneous across electrodes within the same subject, but recovery functions are more variable across electrodes for conditions B and C.

Conclusion
These results suggest that the response of smaller groups of neurons (presumably elicited by the use of a bipolar probe) recovers more slowly and nonmonotonically than the recovery of the aggregate response of the larger group of neurons excited by monopolar stimulation. Results of planned experiments with tripolar probes will be presented, and implications of these results for signal coding by the electrically stimulated auditory nerve will be discussed.

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Sensitivity of Normally-Hearing And Cochlear-Implanted Children Raised In The US And Taiwan To Pitch-Related Cues

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Background
The ability of children to hear subtle changes in pitch is potentially critical for their acquisition of the native language and yet has been little investigated. For children speaking a tonal language, rapid changes in voice pitch carry linguistic information. We hypothesize that these children might have a finer sensitivity to pitch than English-speaking children. Pitch perception by cochlear-implanted (CI) listeners is notoriously poor. However, children who are implanted very early in life may be able to maximize the benefits of brain plasticity. Our second hypothesis is that these children might overcome to some extent their device limitations and process purely temporal pitch more effectively than their normal-hearing (NH) peers.

Methods
Participants were NH children and CI children tested in the US and Taiwan. Percentage correct was measured in a 3-interval 3-alternative forced choice task, for the discrimination of fundamental frequency (F0) of broadband sine-phase harmonic complexes, and for the discrimination of sinusoidal amplitude modulation rate (AMR) of broadband noise. The reference F0 was 100 and 200 Hz, as was the reference AMR. Stimuli were presented via loudspeakers and CI children listened through their speech processors. Data were fitted using a maximum-likelihood technique that extracted threshold and slope.

Results
Preliminary results indicate that in all four groups of listeners, there were large individual differences in threshold and slope, but they did not vary systematically with age. In the NH population, Taiwanese children had lower thresholds and steeper slopes than American children for F0 discrimination but not for AMR discrimination. In the CI population, there was no obvious difference in threshold or slope between Taiwanese and American children.

Conclusion
Based on these preliminary results, our first hypothesis is validated. Speakers of a tonal language seem to display a finer sensitivity to subtle changes in F0. This benefit however does not transfer to purely temporal pitch. Our second hypothesis is rejected: whether one considers the child’s age, the implantation age or the age of use, threshold does not significantly correlate with age. Children who were implanted earlier did not display a better sensitivity to F0 or AMR than children who were implanted later in life. [Work supported by NIH/NIDCD grant no. R21 DC011905 to MC]

Cochlear implant use causes changes in the cochleotopic organisation of auditory cortex in deaf animals

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Background
Neonatal deafness results in the loss of the normal cochleotopic organisation of primary auditory cortex (AI). Environmentally-derived chronic electrical stimulation, via a cochlear implant (CI), can restore that organisation (Fallon et al. 2009, J Comp Neurol). This study explores the time course of the changes in cochleotopic organisation by using chronic recordings of neural activity in auditory cortex of both normal hearing (NH, n=2) and neonatally deafened cats receiving auditory input via a CI (DCI, n=2).

Methods
All animals had 6x10 ‘Utah’ planar silicon substrate recording electrode arrays (Blackrock Microsystems) implanted into putative AI as young adults. Multi-unit cortical responses to acoustic or electric stimulation were obtained under ketamine-xylazine anaesthesia at least monthly for approximately six months.

Results
In the NH animals, well characterised acoustic response areas were initially obtained on 67% of the recording channels, which decreased to 52% in the final recording session. In contrast, in DCI animals, well characterised electrical response areas were initially obtained on 22% of the recording channels, which increased to 69% of channels in the final recording.
session. There was no significant change in cochleotopic (tonotopic) organisation in NH animals over the 6-month recording period (paired t-test on characteristic frequency; p's > 0.1). However, DCI animals exhibited a significant change in cochleotopic organisation over the 6-month recording period (paired t-tests on best electrode; p's < 0.01). The first statistically significant changes in cortical organisation (paired t-tests; p < 0.05) occurred after 4-5 months.

Conclusion
These results demonstrate that auditory cortex can undergo changes in cochleotopic organisation (over several months) as a result of CI use. They also suggest chronic CI use increases cortical responsiveness.

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Bilateral Effects of chronic unilateral Intra-Cochlear Electrical Stimulation on the Central Auditory Pathway
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Background
Little is known about binaural interactions induced by simultaneous acoustic and electric stimulation of the auditory system in patients with unilateral deafness. Since large differences are known between the loudness coding in electric and acoustic hearing, plastic changes would be expected in structures which receive acoustic and electric input in binaural hearing. This study therefore aims to investigate the structural consequences of unilateral chronic intra-cochlear electrical stimulation within the ascending auditory pathway in an animal model.

Methods
Normal-hearing guinea pigs were single-side deafened by the insertion of a standard HiRes 90k® cochlear implant with a HiFocus1j electrode array into the first turn of the cochlea. Four to five electrode contacts were used for the stimulation. Six weeks after surgery, the speech processor (Auria®) was programmed based on tNRI-values and mounted on the animal's back. The three experimental groups were stimulated with a HiRes strategy based on different stimulation rates. Group 1: lowest feasible stimulation rate (275 imp/s). Group 2: medium stimulation rate (1500 imp/s), group 3: highest feasible stimulation rate (5000 imp/s). The experimental groups were compared to group 4: unilateral deafened control (implanted but not stimulated). Experimental groups and controls experienced a standardised free field auditory environment (16 h/day). After 90 Days of stimulation cell density was determined in the medial geniculate body and in the inferior colliculus.

Results
Group 1 (275 imp/s) showed a significantly higher cell density bilaterally in the inferior colliculus (IC) and on the normal hearing side of the medial geniculate body (MGB) compared to the unilateral deaf control. Group 2 (1500 imp/s) showed a significantly higher cell density bilaterally in the MGB and the IC compared to the unilateral deaf control. Group 3 (5000 imp/s) showed a significantly higher cell density bilaterally in the MGB and on the normal hearing side of the IC compared to the unilateral deaf control.

Conclusion
The data of the present study shows that electrical intra-cochlear stimulation has a bilateral and rate independent conservational effect on key structures of the auditory midbrain. The decreased cell density in the control group seems to be induced by the sensory deprivation, which was prevented or reduced by the electrical stimulation of auditory nerve fibres. Based on the present data, an early implantation could be recommended in single sided deaf patients.
Growth-Function of Electrically Evoked Brainstem Responses in Cochlear Implant Patients
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Background
Objective methods are not only important for the fitting process of cochlear implants (CIs) e.g. in babies but can also provide insight about the response of the auditory nerve. Here we measure electrically evoked brainstem responses (eBERA) with single pulse stimuli, study their amplitude-growth function and relate them to the perceived loudness.

Methods
This study was conducted with CI patients with monopolar electrode configuration. Implants were controlled with a research interface box (RIB II). Biosignals were recorded with three electrodes on vertex and neck to measure eBERA responses (ground: contralateral cheek), amplified and digitized. Patients scored loudness with a stylus on a tablet. The tablet was covered with a form which showed five coarse loudness categories but read out was conducted with high resolution.

Results
Because of the huge artifact of the electrical stimulus, the first 1.25 ms of the responses were deleted completely, then responses to anodic and cathodic pulses were subtracted. The remaining stimulation artifact was eliminated by fitting and subtracting an exponential decay function. We then analyzed the amplitude of the wave V peak and identified significant amplitudes with the binominal average method at a confidence level of 99.7%. eBERA amplitudes increased linearly with the stimulation current. We excluded cases with muscle artifacts at high currents, which were probably caused by stimulation of the facial nerve. The figure (see supporting files) show the relation between stimulation strength, wave V amplitude and loudness. Results show significant eBERA responses already at very small loudness categories. The high correlation coefficient indicates a close relation between eBERA amplitude and loudness.

Conclusion
eBERA measurements require stringent data quality evaluations to exclude artifacts. Doing so, we were able to measure significant (99.7%) eBERA responses almost down to perception threshold. eBERA amplitudes grew linearly with stimulus current and also the relationship between loudness and current was well fitted with a linear model. Therefore also loudness grew linearly with eBERA amplitudes. This finding indicates that perceived loudness - at least for single pulse stimulation - is proportional to the number of excited nerve fibers.

Acknowledgement
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Whole-Nerve Responses to Pulse Trains in the Intact and the Degenerating Auditory Nerve in Guinea Pigs

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Background
Effective functioning of cochlear implants (CIs) depends on the integrity of the auditory nerve, and its ability to transduce electrical signals to the central nervous system. Traditional measures such as threshold and amplitude of the electrically evoked compound action potential (eCAP) can be indicative of the nerve’s viability. However, the information that is conveyed by CIs ultimately lies in patterns of pulse trains rather than in its constituting individual pulses. Therefore, by means of eCAP recording, we have characterized the auditory nerve’s behavior under various multiple-pulse conditions.

Methods
Guinea pigs were deafened by co-administration of kanamycin (400 mg/kg) and furosemide (100 mg/kg) two or six weeks before acute experiments. Untreated guinea pigs were used as normal-hearing controls. eCAPs were recorded using a MED-EL PULSAR cochlear implant. Pulse trains consisted of alternating biphasic current pulses, and had a duration of 10 or 100 ms; pulse rate was increased from 62.5 Hz to 2.5 kHz in ten steps. Amplitude and latency of the last ten eCAPs in the pulse train were measured. Immediately after the acute experiments, the animals were sacrificed for histological analysis of the spiral ganglion cells.

Results
The eCAP amplitude to the first pulse of the pulse train decreased with duration of deafness, which correlates with the decrease in number of spiral ganglion cells. The eCAP amplitude to later pulses decreased with higher pulse rates. This decay was more pronounced with longer pulse train duration, and also more pronounced for normal-hearing animals. In each animal we observed an oscillating effect in eCAP amplitude for pulse rates around 1.5 kHz. The average oscillating frequency was between 600 and 800 Hz. The amplitude of this oscillation was significantly smaller for normal-hearing than for deafened animals. Finally, the eCAP latency increased with increasing pulse rate, but only significantly so for the normal-hearing animals.

Conclusion
The slower decay and shorter latency of eCAPs in deafened animals may be a result of degeneration of the organ of Corti, creating a simpler neural interface. Higher oscillation amplitudes in these animals could indicate increased synchrony of the population of nerve fibers.
Recognition of Complex Frequency-Modulation Patterns Based on Recovered Envelope Cues in Cochlear Implant Users: Effects of Modulation Depth and Peripheral Filtering

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Background
Frequency-modulation (FM) patterns convey crucial information for sound identification. FM is encoded into both temporal-envelope and temporal fine-structure cues (TFS) via cochlear filtering and neural phase locking in the auditory periphery. Cochlear implant (CI) users provide a unique opportunity to assess the ability to use the envelope cues reconstructed from FM after peripheral filtering with minimal contribution of TFS. This study also aims at assessing the effect of the clinical processor on the transmission of FM via the envelope reconstruction mechanism.

Methods
The sensitivity of CI users and normal-hearing (NH) listeners was measured in a discrimination task using complex FM patterns applied to a 930-Hz sine carrier. Two factors assumed to affect envelope reconstruction were tested: (i) the extent of frequency excursion in FM stimuli, and (ii) the characteristics of peripheral filtering. Three sets of FM stimuli were created using a low, medium or high modulation index. The FM stimuli were then passed either through a model of the normal cochlear filter (gammatone) or the corresponding broader clinical filter of the CI. In each case, the stimuli were presented through one electrode of the CI at the same level.

Results
For most conditions, CI users had better sensitivity to FM patterns on the basis of reconstructed envelopes than NH listeners. This might result from the use of a linear input-output function for the CI processor. Sensitivity increased substantially with frequency excursion and the effect was larger for CI users. However, CI users were not able to benefit from the use of filters narrower than clinical filters to recover envelope cues: only NH listeners showed better sensitivity with the gammatone filter compared to the clinical filter.

Conclusion
The results indicate that discrimination of complex FM patterns can be achieved on the sole basis of reconstructed envelope cues. The fact that peripheral filtering, which simulated normal cochlear filtering, provided no benefit over standard clinical filters suggests that current clinical filters are sufficiently selective to promote efficient envelope reconstruction.
Cortical Hemodynamic Response to Speech Stimuli in Adult Cochlear Implant Users by Functional Near-Infrared Spectroscopy

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Background
Cochlear implants (CIs) are a therapeutic option to restore hearing in people with profound sensorineural hearing loss. Objective measures of a CI’s function are needed, especially in the pediatric population where optimal CI programming is essential for normal speech and language development and patient feedback during programming is more difficult to obtain. Functional near-infrared spectroscopy (fNIRS) is a non-invasive, CI-compatible imaging method used to monitor cortical hemodynamic changes. In our previous work using an instrument with 2 light sources and 4 detectors, we showed that acoustic speech stimuli elicit cortical responses in adults and children with normal hearing, as well as pediatric CI users. With this project, we aimed to demonstrate that a response to acoustic speech stimuli by auditory cortex is present with the CI turned on, but not with the CI turned off.

Methods
We collected fNIRS data using 16 light sources and 24 detectors to permit spatial localization of the cortical responses. We tested this approach by using fNIRS to assess for hemodynamic responses in the auditory cortex to acoustic speech stimuli in adult CI users both with their device turned on and with it turned off.

Results
Most participants demonstrated speech-evoked cortical activity only when their CI was turned on and not when it was turned off, as well as substantially larger cortical volume of activation when their CI was turned on compared to when it was turned off. A bilateral cortical response was the most commonly observed finding, regardless of the side of implantation. Participants who showed equivalent cortical responses and volumes of activation with their CI active and inactive were those with recently activated CIs.

Conclusion
Our findings show that fNIRS is a valid tool for measuring cortical hemodynamic responses in adult CI users and hence assessing CI functionality. This method holds potential to provide an objective measure of speech perception among CI users, and may be useful during CI programming.
Electrical Stimulation of the Cochlear Nucleus: Effects of Location and Pulse Rate on Inferior Colliculus Responses

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Background
Auditory brainstem implants (ABIs) help provide some hearing to patients with non-functional auditory nerves or ossified cochleae. Yet, performance of ABI users in speech reception tasks is usually worse than for cochlear implant users. ABIs bypass the cochlea and auditory nerve and electrically stimulate the surface of the cochlear nuclear complex. Two of the possible reasons for their limited performance are: (1) the relative imprecision of device placement, making it difficult to specifically target frequency regions important for speech understanding, and (2) the poor spatial selectivity of electrical stimulation, which limits frequency resolution. Our goal is to evaluate the effects of electrode placement and pulse rate on spatial selectivity and stimulation threshold in the hope of improving ABI performance.

Methods
The surface of the dorsal cochlear nucleus (DCN) of anesthetized rats was stimulated with either commercially available parylene-insulated platinum-iridium twisted electrodes or with novel, flexible micro-electrode arrays integrated into a polyimide polymer pad. Multiunit activity was recorded with a 16-micro-electrode array in the inferior colliculus (IC).

Results
Consistent with the tonotopic organization of the DCN and IC, we found that DCN surface stimulation laterally (low frequencies) to medially (high frequencies) led to lower threshold activation dorsally to ventrally in the IC. Spatial selectivity of electrical stimulation, although somewhat narrower in the rostral-most or caudal-most portions of the DCN, was usually poor. Lower stimulation thresholds were found on the medial side of the DCN. Pulse rate (11 – 513 Hz) did not have a significant effect on these measures across animals.

Conclusion
Our results point to the need for new stimulation strategies and devices to improve frequency representation in the auditory system of ABI users. Ongoing work in our laboratory is investigating whether electrode arrays that conform to the surface of the CN may help in this regard.
Are Amplitude-Dependent Changes in Temporal Resolution in Deaf Cat Inferior Colliculus Affected by Electric Hearing History?

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Background
In previous studies we have reported that temporal resolution of central auditory neurons can be altered by deafness and by experience with intracochlear electric stimulation (ICES). In deaf animals, degradations in the ability of neurons to follow repetitive electric signals and response latencies can be improved by chronic ICES or even restored to normal levels by behavioral training. The present study investigates whether temporal processing in animals with different deafness histories and electric hearing experience is differentially affected by the electric stimulus amplitude used in the electrophysiological experiments.

Methods
Kittens were neonatally deafened by injection of ototoxic drugs. Animals were implanted unilaterally with a feline prosthesis either as juveniles or as adults after long-term deafness (≥3.5 yr), and a regimen of continuous ICES was initiated (~4 h/day, 5 day/wk). Animals were studied after several weeks to months of ICES. Acutely deafened adult animals served as controls. Pulse trains (~10 to 300 pps) were presented at stimulus amplitudes between 2 and 6 dB above neuronal threshold. Neuronal responses were recorded in the external (ICX) and central nucleus (ICC) of the inferior colliculus. Temporal response parameters of isolated single units were quantitatively evaluated, including: 1) maximum stimulus rates that neurons followed in a phase-locked manner (Fmax) and 2) minimum neuronal response latencies.

Results
In all groups, increases in stimulus amplitude resulted in a significant decrease in response latencies in both ICX and ICC neurons. With respect to Fmax, only ICC neurons in juvenile animals showed a significant increase in frequency following with increasing stimulus amplitude. Otherwise increases in stimulus amplitude had no effect on frequency following of neurons in either the ICX or the ICC.

Conclusion
Electric hearing history had no differential effect on amplitude-dependent changes in response latencies in the ICX or the ICC. With the exception of ICC neurons in juvenile animals, hearing history, likewise, had no differential effect on amplitude-dependent changes in Fmax.

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Temporal responses to regular-interval and alternating-interval electric pulse trains in the inferior colliculus

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Background
With cochlear implants, the fundamental frequency and other temporal aspects of a sound can be conveyed by the temporal pattern of evoked neural activity. In previous work, Carlyon et al (2008) used acoustic click trains, bandpass filtered to remove “place of excitation” cues, in order to simulate several aspects of a “purely temporal” pitch code. For example, a click train in which the inter-pulse intervals alternated between 4 and 6 ms produced a pitch equal to that of a regular-interval (isochronous) train with IPI=5.7 ms. A similar result was observed for cochlear implant listeners when presented with a train of biphasic electrical pulses. Additionally, in a guinea pig model, Carlyon and colleagues measured smaller compound action potential (CAP) amplitudes following the shorter interval of a “4-6” acoustic click train, suggesting a potential neural basis of the observed perceptual effect. To obtain a more complete account of the neural representation of pitch in CI users we have recorded the responses of single units from the inferior colliculus of the guinea pig to similar electrical pulse trains.

Methods
In urethane-anesthetized adult guinea pigs, the inferior colliculus (IC) was exposed following a craniotomy and aspiration of the overlying cortex. A tungsten microelectrode was lowered into the brainstem, and responses to acoustic tones were obtained to verify placement. After deafening with neomycin, a PtIr ball electrode was secured inside the scala tympani. A
series of single pulse and pulse train stimuli were presented in blocks at several current intensities. Unit data were processed offline to remove electrical artifacts, classify action potentials, and analyze spike timings.

Results
The majority of units had sustained or build-up temporal response patterns at current levels within ~10 dB of threshold, and 85% of these responded synchronously to the 10 ms period of the 4-6 alternating-interval pulse trains. The remaining units responded only to the onset or offset of the trains. At relatively low current levels, the spiking probability of synchronized units following the 4 ms intervals was usually smaller than that following the 6 ms intervals, similar to the CAP results. At higher levels, however, this difference was less pronounced.

Conclusion
In general, the diverse response patterns to alternating-interval pulse trains were reflected in the responses to isochronous pulse trains and pulse pairs presented at corresponding inter-pulse intervals. How these temporal response patterns relate to pitch perception remains to be elucidated.

Completely Implantable Artificial Organ of Corti
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Background
Present commercial cochlear implants are limited by power consumption, latency, and physical unwieldiness. Efforts to develop smaller and more efficient implants have focused on the use of microelectromechanical systems (MEMS) technology to build small, passive, frequency-filtering artificial basilar membranes (ABMs) to replace signal processing. We present a piezoelectric cantilever array that is designed to replace the function of the organ of Corti. Fluid motion in the ST deflects an array of piezoelectric cantilevers; this deflection produces an electric potential on the outer electrodes of each cantilever that is transmitted into the surrounding perilymph. When implanted, this device would function as a low power, completely implantable artificial organ of Corti.

Methods
An implant was fabricated using batch MEMS processing techniques. A silicon backbone supports five 400µm wide, 5µm thick bimorph (Platinum/aluminum nitride (AlN) stack) cantilevers, each of which is designed to have a resonance corresponding to its tonotopic location in the guinea pig scala tympani (ST) (20-40kHz). A 1mm wide, 10µm thick parylene and gold ribbon cable extends 3cm from the probe to an electrode bay, where electrical connections to each cantilever are accessed. Actuation responses of the fabricated devices were measured using laser vibrometry.

Results
A model predicting the resonant behavior of cantilevers in a viscous fluid was used to determine the geometry of the array. Further modeling predicts the current output of the device in the guinea pig cochlea ST; this model predicts the device will produce 90dB SPL ABR thresholds when actuated by a 1Pa sound source. Fabrication challenges such as film stress passivation and selective etching have been identified and the next generation prototypes are designed to overcome them. First generation devices have been fabricated and are electrically addressable through the ribbon cable and a MEMS electrode bay (developed to monitor/control the output). Individually actuated cantilevers show good agreement with model predictions. In vivo implantation of inactive devices in the guinea pig for a week produced no infection but some granulation tissue surrounding the device.

Conclusion
A novel, completely implantable artificial organ of Corti was designed to passively restore hearing using a piezoelectric cantilever array. Initial in vitro electromechanical array element testing shows good agreement between predicted and measured actuated cantilever frequency response. Future work includes in vitro device response testing in water and ionic fluid to fully characterize the array behavior, as well as in vivo testing to measure ABR thresholds.
Correlation between intracranial pressure and skull vibration in bone conduction of cadaveric human whole heads

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Background
Several pathways including ear canal radiation, inertia of the ossicles and cochlear fluid, compression and expansion of the skull, and pressure transmission through soft tissues are considered to contribute to bone conduction hearing. The importance and frequency dependency of these pathways have not been comprehensively understood. Especially, the contribution of pathway via non-osseous skull contents such as brain content and CSF are controversially discussed. The aim of this study is to assess the correlation between intracranial pressure and skull vibration on the promontory for bone conduction stimulation.

Methods
The measurements were performed on 3 human cadaver heads. The bone conductive stimuli of 100 dB SPL in the frequency range of 0.1 – 10 kHz were delivered to the cadaver heads via bone anchored hearing aid (BAHA). The BAHA was held in place with a 5-N steel headband at mastoid, forehead, eye, and neck. Additionally, it was percutaneously implanted at the mastoid and forehead. With the bone conductive stimulation, the motion of the cochlear promontory and intracranial pressure at the center of the head were measured simultaneously using a Laser Doppler Vibrometer and a hydrophone. An absolute static pressure sensor placed in the middle of the head during the measurement to monitor intracranial static pressure.

Results
Frequency dependency of the intracranial sound pressure differed from the corresponding dependency of the skull bone vibration measured at the cochlear promontory on the ipsilateral and contralateral sides. Further, the intracranial attenuation could be estimated from the difference of skull bone vibration between the ipsilateral and contralateral sides.

Conclusion
The presence of variation of intracranial sound pressure indicates that non-osseous skull contents may contribute to bone conduction hearing in some frequencies.
Functional and morphological effects of variable laser stimulation parameters at the ear drum level

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Background

Visible light can be used to activate the peripheral hearing organ when applied at the ear drum level (Wenzel et al. 2010). However, to enable the detection of complex sounds and intelligible speech perception, controlled frequency-specific activation spanning the audible frequency range is necessary. Using current laser technology, it is not possible to switch effectively between different laser sources, with different wave lengths, within the duration of a word. Therefore, the overall objective of our project is to establish methods for achieving frequency specific activation of the complete audible spectrum using monochrome laser pulses. As a first step, we analysed the electrophysiological and morphological effects of varying laser-pulse repetition rates in an animal model.

Methods

Click-Auditory Brainstem Responses (ABR) were recorded preoperatively in anesthetized guinea pigs to confirm normal hearing. A 100 µm diameter optic fiber was inserted into the outer ear canal and directed towards the ear drum. Optically-induced ABRs were recorded in response to laser stimulation with Q-switched 10 ns, 532 nm laser pulses from a diode-pumped solid state Nd:YAG laser (Xiton Photonics GmbH, Kaiserslautern, Germany). The animals were then sacrificed immediately after laser irradiation and the temporal bones were rapidly explanted and processed for histological analysis.

Results

Optical stimulation of the ear was possible with pulse energies from 0-17 µJ/pulse and repetition rates from 10 Hz - 20 kHz. Increasing the laser-repetition rate led to a superposition of the optically-induced ABRs. The morphological changes were dependent on the applied energy level, time of exposure, and pulse repetition rate.

Conclusion

These findings suggest that laser-pulse shaping based on variable temporal pulse distributions, a process that is well known in physics, applies for biological systems as well. Further studies investigating optimal pulse shaping strategies for the activation of the complete auditory spectrum as well as the biocompatibility of green light irradiation of the auditory system are under investigation.

Fabrication of biomimetic shaped ceramic Ossicles and Evaluation of their Vibration Performance using Finite Element Method

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Background

Many three-dimensional geometric models of the human ossicles have been proposed and finite element modeling has distinct advantages in modeling the middle ear over circuit models that correspond indirectly with its structure and geometry. The object of this study was to evaluate vibration performance of synthetic ossicular chain fabricated by projection-based microstereolithography (pMSTL) and to compare its vibration characteristics with those of human ossicles. We also investigated the characteristics of the fabricated ossicular chain using finite element analysis method.

Methods

Synthetic ossicular scaffolds, which have same configuration of human ossicular chain, were created from hydroxyapatite, porogen and SK-cytec using pMSTL and CAD/CAM technology. Their vibration performances resulting from piezo-electric and signal analyzer were measured with laser vibrometer. Vibration characteristics of the ossicles were also measured by finite element analysis.

Results

Fabricated ossicles showed output resonance frequency of 528, 1344, 1921, 3584, 5505, 6060, 7168, 7894, 11488 and 17302 Hz with input resonance frequency of 5520 and 6849 Hz, which was comparable to known results of human
Conclusion
Fabricated ossicular chain using pMSTL and CAD/CAM technology may have similar vibration characteristics with those of human ossicular chain.

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Design and Preliminary Testing of a Visually-Guided Hearing Aid
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Background
The defining complaint of listeners with sensorineural hearing loss (SNHL) is difficulty hearing in "noise." For most people with SNHL, hearing aids are the only effective treatment. Despite the many advances in hearing aid design over the past few decades the fundamental problem remains: how to selectively amplify the sounds the listener wishes to hear while excluding unwanted, interfering sounds. The goal of this work is to explore a new approach to amplification designed to facilitate sound source segregation and selection. This approach is based on the premise that only the listener can make the distinction between which sources deserve attention and which should be ignored, and therefore which sources should be amplified and which suppressed. In this study we describe initial efforts at designing a "visually-guided hearing aid" (VGHA) that senses where vision is directed and steers a beam of amplification in that direction.

Methods
The new prototype "hearing aid" couples an acoustic beam-forming microphone array to an eye-glasses-mounted eye-tracking device. The array consists of four pairs of cardioid microphones secured on the headband of a pair of earphones. The gaze angle sensed from the eye tracker is used to steer the acoustic look direction (ALD) of the array. Estimates of the benefit of the VGHA include acoustic measurements of array gain/attenuation as a function of azimuth surrounding the ALD and perceptual measurements of speech recognition under multitalker conditions. Furthermore, estimates were obtained of the time required for dynamic refocusing of the ALD during speech testing.

Results
Acoustic measurements conducted on the array in a mildly reverberant test environment indicate spatial tuning that varied with frequency but that provided a high degree of selectivity at frequencies above 2 kHz. Perceptual measurements indicated that near-normal selectivity of a target talker may be achieved by the VGHA, as reflected in the magnitude of spatial release from speech-on-speech masking. The interface and components working together were able to reposition the ALD rapidly across a range of azimuths spanning roughly ± 45°.

Conclusion
The approach embodied by the VGHA offers some advantages over current methods for amplification and sound source selection in uncertain and dynamic multitalker listening environments. The current prototype only tests the premise upon which the device depends and by no means comprises a wearable, useable device. However, as a platform for testing this new approach the VGHA appears to be useful.
Assessment of temporal resolution of bone-conducted ultrasonic hearing using psychophysical and neuromagnetic measurements

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Background
Bone-conducted ultrasound (BCU) is perceived even by the profoundly sensorineural deaf. We have objectively proven the BCU perception by neurophysiological data and developed a novel hearing-aid using the BCU perception (BCU hearing aid: BCUHA) for the profoundly deaf, that transmits a 30-kHz bone-conducted carrier that is amplitude-modulated by speech or environmental sounds. In this study, the temporal resolution of the BCUHA hearing were assessed by psychophysical and neuromagnetic measurements in normal-hearing and profoundly deaf subjects.

Methods
To investigate the temporal resolution of the BCUHA hearing psychophysically, the temporal transfer modulation functions (TMTFs) were measured. TMTF shows the threshold of sinusoidal amplitude-modulation (SAM) detection as a function of modulation frequencies. The SAM detection threshold is determined systematically by measuring a detection of modulation depth. TMTF for 10-, 20-, and 30-kHz bone-conducted (BC) sounds were measured. Further, mismatch fields (MMFs) for changes of stimulus-duration were recorded. MMFs for 10-, 20-, and 30-kHz bone-conducted tone bursts amplitude-modulated by 1 kHz, and a 1-kHz air-conducted (AC) tone burst were measured in different sessions. Each session consisted of one standard stimulus (75-ms duration, 85%) and three types of deviant stimuli (52.5-, 37.5-, and 22.5-ms durations, 5 % each).

Results
The psychophysical detection threshold curves for all carriers commonly showed the low-pass characteristics at low modulation frequencies; the thresholds were roughly constant at modulation frequencies from 10 to about 100 Hz and then begin to increase at about 100-150 Hz as the modulation frequency increases. Meanwhile, clear MMFs were observed for all types of stimuli. Equivalent current dipoles (ECDs) moments of MMFs were significantly different between three types of BC and AC stimuli (10-kHz BC:20-kHz BC:30-kHz BC:1-kHz AC=0.95:0.92:0.91:1.0), however, ECD moments of BC stimuli were close to that of AC stimulus.

Conclusion
The both results obtained in the psychophysical and neuromagnetic measurements indicated that the BCUHA hearing has nearly equal temporal resolution to that of lower-frequency BC and AC sounds. These results also provide useful information for further developments of the BCUHA.
Development of a Bedside Gaze Stabilization Test
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Background
The gaze stabilization test (GST) is used not only in examining gaze stability required by daily life activities such as walking, running, and driving, but also in identifying unilateral or bilateral vestibular deficits. However, a computerized GST (CGST) is an expensive assessment which cannot be commonly used in most balance and vestibular clinics. Validation of low-cost and low-technical clinical “bedside” tests is required to decrease health care costs.

Methods
All twenty healthy volunteers were between the ages of 20-60 ears and had normal extra-ocular range-of-motion. This study measured the effect of a bedside GST (BGST) as a validated screening tool to accurately identify adults with normal inner ear balance function, as well as the test-retest reliability of BGST when re-assessed 5-7 days after the initial testing session. Subjects were instructed to identify a visual target placed at 0.2 logMAR increased from static visual acuity score while actively and passively performing head movements in the yaw plane at an initial screening velocity of 130 degs/sec.

Results
The main outcome was a strong positive correlation in both active and passive BGST head movement amplitude degree and head movement velocity. Our finding showed that passive head movements had good specificity (85%) to identify healthy individuals with functioning peripheral vestibular systems. The BGST seems to be a reliable screening assessment to identify normal functional performance of the peripheral vestibular system to maintain stable gaze while moving the head at a screening velocity of 130 degs/sec and has good test-retest reliability in head movement amplitude degree (r = 0.795) and head movement velocity (r = 0.797) of passive BGST within two testing sessions.

Conclusion
A low-cost, low-tech (bedside) version of the GST may not only play an essential part in reducing the burden of medical expenses for adults but may also be useful as an assessment tool to screen individuals with vestibular impairment. In this study, the BGST screening assessment was deemed to be a measure to identify individuals with normal reported peripheral vestibular function and it showed good test-retest reliability in degree and velocity of passive BGST. The aim of this future research is to establish a screening cut-point score in terms of the maximum head velocity that will identify individuals with both normal peripheral vestibular function and peripheral vestibular impairment.
Masking effects in patients with auditory neuropathy - possible involvement of suppression mechanism caused by normal outer hair cell function -
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Background
Masking is a phenomenon by which the presentation of an interfering sound (masker) decreases a signal’s audibility. Noise audiometry is a proposed method for assessing hearing acuity in noisy environments based on measurement of auditory thresholds in the presence of masking noise. Auditory neuropathy is a type of sensorineural hearing loss characterized by absence or marked abnormality of auditory brainstem responses beyond that expected for the degree of hearing loss, with preserved activity of the outer hair cells. Noise audiometry has not been examined extensively in patients with auditory neuropathy, although speech intelligibility in patients with retrocochlear lesion often deteriorates remarkably in the presence of masking noise. To investigate the possible mechanism of the characteristic masking feature of the disease, this study reviewed the findings of noise audiometry and pathogenesis retrospectively in patients with auditory neuropathy.

Methods
Study Design: Retrospective chart review.
Patients: 55 ears of 30 patients with sensorineural hearing loss who underwent noise audiometry at our institute since 2010 in response to complaints of hearing difficulty in noisy environments.
Main Outcome Measures: Masked threshold for narrow band and white noise.

Results
Masking effects in patients with auditory neuropathy were significantly greater than in patients with hearing loss of other types. Specifically, masking effects of broad-band white noise were greater than those of narrow-band noise. With white noise, masking effects were observed even in the elevated threshold region, where little contribution of excitatory masking effect is expected.

Conclusion
This retrospective study of noise audiometry conducted at our institute revealed strong masking effects, especially in patients with auditory neuropathy. Fragility for the masker in patients with retrocochlear lesion is often explained as “easy fatigability” or “easy adaptability”, known as a temporal threshold shift in Bekesy audiometry. However, such an explanation might be inappropriate to explain the masking phenomenon caused by the lower level masker, which presumably does not cause excitatory masking effects as a line busy or an adaptation mechanism. We speculate that other masking effects originating from cochlear nonlinearity caused by outer hair cells, such as “suppression mechanism”, are involved in masking by a lower level broad-band masker because auditory neuropathy type hearing loss exhibits normal outer hair cell function.
Clinical features and prognosis of hearing impairment due to meningeal carcinomatosis

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Background
Meningeal carcinomatosis is uncommon and characterized by multifocal spread of neoplastic cells in the leptomeninges. A variety of multiple neurological symptoms and known history of malignancy are usually the signs suggesting the diagnosis. Only a few reports of hearing impairment due to meningeal carcinomatosis have been reported. By reason of their poor vital prognosis, hearing impairment has not been evaluated and discussed enough. Here we presented four cases of progressive hearing impairment due to meningeal carcinomatosis.

Methods
We conducted a retrospective chart analysis of patients demographic, audiograms, and MRI findings from case series presenting hearing impairment due to meningeal carcinomatosis. Auditory brainstem response (ABR) and distortion product otoacoustic emission (DPOAE) were also tested.

Results
Primary lesions were two lung adenocarcinoma, one esophageal carcinoma and malignant lymphoma respectively. Diagnosis was made based on past history of malignancy, MRI findings, and the cytology in cerebrospinal fluid. Major complaining symptoms were rapidly progressed hearing impairment, unsteadiness, and tinnitus. Facial nerve palsy was not noted at first, but two cases presented in later stages. All cases showed enhancement of vestibule-cochlear nerves on gadolinium-weighted MRI. Two cases of internal auditory canal metastasis were revealed on high-resolution MRI focused on internal auditory canal. ABR obtained in two cases showed prolonged latency between wave I and III. The response of DPOAE was initially seen in lower frequencies, but finally lost completely. Hearing impairment rapidly progressed to deafness even in the patient with high-dose steroids treatment. Hearing improvement was not observed, except for the case with malignant lymphoma.

Conclusion
Hearing prognosis was poor in hearing impairment due to meningeal carcinomatosis. Steroids treatment had the least effect on preventing the progression of hearing impairment. Internal auditory canal metastasis often coexisted with meningeal carcinomatosis and suggested cochlear dysfunction. Both gadolinium enhancement and high-resolution MRI focused on internal auditory canal were recommended to diagnose the meningeal carcinomatosis as soon as possible. Clinical course and results of ABR and DPOAE suggested that cochlear nerve dysfunction preceded and cochlear dysfunction developed later.

Aided ASSR on a Cadaver: Relevance in the Clinic

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Background
Auditory Steady State Response (ASSR) is used to determine aided hearing thresholds, set hearing aid (HA) parameters and verify HA benefit. Aided auditory evoked potentials (AEPs) can be affected by electromagnetic interferences related to the HA (Crose et al, 2011). Most studies on aided ASSR, use an electrode montage with reference electrode physically distant from the HA (e.g. Shemesh et al., 2012); however studies (Picton et al., 1998; Stroebel et al., 2007) have used the mastoid ipsilateral to the HA for the reference electrode. Additionally, braiding minimizes electrical interference (Stecker and Patterson, 1996). However, AEP studies, especially ones using devices like HAs, do not mention if electrode leads were braided. The purpose of this study was to investigate if the amplitude spectrum of the ASSR is affected by 1. HA-related interference based on the position of the reference electrode (ipsilateral vs. contralateral); 2. Braiding electrode leads.

Methods
An adult male cadaver was fitted monaurally with an Oticon Safari SuperPower HA to record “pure electrical interference” without interaction with physiological responses. HA specifications are listed in Table1 and the setup is described in Figure1. Recordings were made from the forehead (Fz) to 1. Ipsilateral mastoid; 2. Contralateral mastoid; with clavicle as ground and with electrode leads unbraided vs. braided. Before data collection, we recorded ASSR with the stimulus at 90 dB SPL but with the HA ‘OFF’ and the stimulus at 0 dB SPL but the HA ‘ON’ to ensure that possible interferences were from the HA. Multi-frequency ASSRs were recorded from 90- to 40 dB SPL in 10 dB decrements for each montage-
unbraided and braided. Also, single frequency ASSRs were recorded at 90 dBSPL for each montage- unbraided and braided.

Results
A peak indicating electrical interference was observed in the amplitude spectrum of the multi-frequency ASSR at approximately 120 Hz for the ipsilateral-unbraided condition for all intensities tested and also for the single frequency stimulus, ipsilateral-unbraided condition at 90 dBSPL. Such interference can potentially affect the acquisition of a response clinically. However, this interference disappears for ipsilateral-braided, contralateral-unbraided and contra-braided conditions (Figure2).

Conclusion
Positioning the reference electrode away from the HA and braiding electrode leads is essential while recording AEPs, especially in the presence of a HA. This study is unique in nature because it highlights the possibility of electrical interference from a HA and emphasizes the utilization of simple techniques that can eliminate such interference.
**Specifications: Oticon Safari SuperPower**

<table>
<thead>
<tr>
<th>Ear Simulator</th>
<th>Specifications</th>
</tr>
</thead>
<tbody>
<tr>
<td>OSPL90</td>
<td>Peak 143 dBSPL</td>
</tr>
<tr>
<td>1600 Hz</td>
<td>131 dBSPL</td>
</tr>
<tr>
<td>Average</td>
<td>136 dBSPL</td>
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<tr>
<td>Full-on gain</td>
<td>Peak 82 dB</td>
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<tr>
<td>1600 Hz</td>
<td>73 dB</td>
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<td>Average</td>
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<tr>
<td>Frequency Range</td>
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<td>Total Harmonic Distortion</td>
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</tr>
<tr>
<td>(input: 70 dBSPL)</td>
<td>800 Hz 1.0%</td>
</tr>
<tr>
<td></td>
<td>1600 Hz 2.0%</td>
</tr>
<tr>
<td>Equivalent Input Noise Level</td>
<td>Omni 20 dBSPL</td>
</tr>
<tr>
<td></td>
<td>Dir 31 dBSPL</td>
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Table 1: The Oticon Safari SuperPower hearing aid was used in the present study. The hearing aid specifications are listed in the table above.

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**Effect of Threshold Sound Conditioning on Pure-Tone Hearing Thresholds in Human Subjects with Sensorineural Hearing Loss**

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**Background**

Sensorineural hearing loss has long been considered to be an irreversible phenomenon. Most likely for this reason, there has been little or no attempt to investigate the effect of sound conditioning on the hearing threshold in people with sensorineural hearing loss. Because we have observed the same phenomenon (i.e., a hearing threshold decrease due to sound conditioning) during research on tinnitus treatment, we decided to investigate the effect of sound conditioning on pure-tone hearing threshold in people with sensorineural hearing loss. Sound conditioning using a non-traumatic,
moderate-level acoustic signal is a well-studied method to protect against age-related or noise-induced hearing loss in animals. In humans, acoustic signals provided at the hearing threshold level or slightly higher than the hearing threshold level have been studied for the treatment of hyperacusis and tinnitus and have been found to have positive effects. In this study, we adopted the term “sound conditioning” instead of “sound training” because the subjects were not asked to pay attention to the acoustic signals. In addition, because the subjects in the experimental group were instructed to listen to the acoustic signals at their hearing threshold levels (i.e., a barely audible level), we named the experimental method “threshold sound conditioning (TSC).”

Methods
We examined the changes in the pure-tone hearing threshold after 2~3 weeks of TSC (i.e., sound conditioning with a frequency-specific acoustic signal at the hearing threshold level) or music listening (i.e., listening to music at a comfortable listening level). In addition, we examined hearing threshold changes without TSC or music listening in both groups.

Results
After the music listening period in the control group, the only change observed was a hearing threshold increase at one frequency. In contrast, statistically significant decreases in the hearing threshold were observed at several frequencies after the TSC period. Clinically significant threshold decreases of 10 dB or more were observed at 9 subjects in the experimental group. In total, clinically significant threshold decreases were observed for 33 frequencies. The average hearing threshold of the 20 TSC frequencies before the break period, after the break period, and after the TSC period was analyzed. A hearing threshold decrease of 12.6 dB was observed for the TSC frequencies of the TSC ear (P < 0.001).

Conclusion
Threshold sound conditioning can decrease pure-tone hearing thresholds in people with sensorineural hearing loss.
Using the Cochlear Microphonic as a Clinical Diagnostic Procedure to Detect the Cochlear Location of Damage of Outer Hair Cells
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Background
The progression of new medical treatments for curing sensorineural hearing loss requires the development of new diagnostic procedures that localize the damaged site/s within the cochlea, and/or auditory nerve. The goal of our research is to investigate the cochlear microphonic (CM) as a clinical diagnostic procedure to detect the location and amount of damage in hearing impaired ears.

The CM is an electrophysiological signal generated from receptor currents passing through outer hair cells (OHCs). We have recently shown that CM amplitude, in response to a low frequency tone embedded in filtered noise, increases as the high-frequency cutoff decreases (Chertoff, Earl, Diaz, and Sorensen, (in press)). When the CM amplitude is normalized relative to the unmasked response and plotted as a function of cochlear distance from the apex, the growth is defined as the cumulative amplitude function (CAF).

Methods
We have studied the CAFs in 6-week-old Mongolian gerbil by recording CM responses to a 762 Hz tone from 80 to 40 dB SPL levels embedded in 25 high-pass filtered noise conditions.

Results
We observed that the CAFs reached an asymptote at distances more apically than the CAF for high level signals. This suggests an apical shift in the contribution of OHCs to the CAF. An important question is the contribution of signals coming from other cochlear structures to the CAF. That is, at low frequencies, phase-locked action potentials may also contribute to the electrophysiological cochlear response.

Conclusion
Thus determining the contribution of neural potentials to the CAF will be important for determining the specificity of the CAF to OHCs.

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Steps towards untangling the enigma of auditory processing disorder
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Background
Listening problems in children, despite apparently normal hearing, are common in a variety of developmental learning disorders. In paediatric audiology, these problems are often diagnosed as ‘auditory processing disorder’ (APD). The validity of APD as a diagnostic category is controversial since many children receiving a diagnosis of APD are, largely, indistinguishable from children with other disorders such as specific language impairment (SLI). It has been argued that the diagnostic label that children receive reflects referral route rather than valid differences in auditory or linguistic abilities. To counter this, we hypothesise that there are clear differences in presenting symptoms and that these determine the referral route a child is most likely to follow.

Testing this hypothesis requires three steps, two of which are addressed here.

Methods
First we developed a psychometrically robust tool for assessing the listening difficulties. Parents of 950 children indicated the extent to which they agreed with 84 possible presenting symptoms. Based on their responses, five factors were extracted: Online speech/nonspeech processing; Environmental sensitivity; Language & literacy; Pragmatic skills; and, Memory. Second, we compared the responses of parents of 3 groups of children commonly thought to have listening difficulties i.e., SLI (n=23); SLI/APD (n=18) and dyslexia (n=19).

Results
The 3 groups of children received significantly different parental evaluations [F(2, 57) 4.58, p =.014], and they could be ordered hierarchically in terms of severity of listening difficulties, from least (dyslexics) to most (SLI/APD) severely affected on the latent traits gauged by this questionnaire. The dyslexics scored in the normal range on all factors except Language & literacy and Memory. The children with SLI/APD had difficulties across all latent traits except Environmental sensitivity.

Conclusion
The questionnaire developed suggests there are measurable differences between the different groups assessed here. In the final step towards answering our primary research question, we will use the questionnaire to compare children clinically referred for assessment for APD, with children clinically referred to speech and language therapists and to determine how and if these two groups of children differ in their presenting symptoms.

Crossed and uncrossed acoustic reflex growth functions in normal hearing children, adults and children with auditory processing difficulties
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Background
Children who experience difficulty understanding sounds and learning through their hearing in spite of normal sensitivity are often considered for evaluation of auditory processing difficulties. Professional associations recommend a battery of tests that assess behavioral function, often in difficult listening situations, and the evaluation of underlying neural integrity of the auditory system using objective measures. But there is little agreement on what objective measures should be used. Acoustic reflex measurements may provide a sensitive objective measure of lower brainstem integrity, particularly when patterns of crossed and uncrossed reflexes are examined. Our previous work has noted a higher incidence of threshold elevations or absent acoustic reflexes in children with suspected Auditory Processing Disorder (APD), especially in the crossed reflex pathway. This study presents a more detailed description of acoustic reflex characteristics in children with suspected APD through the measurement of acoustic reflex growth functions. These functions describe the rate of change in middle ear impedance with changes in stimulus level. Evaluations of both crossed and uncrossed pathways were examined.

Methods
Uncrossed (ipsilateral stimulus and recording) and crossed (contralateral stimulus and recording) reflexes were recorded in normal hearing adults and children, and in children with suspected APD. Stimuli were pure tones of 500, 1000 and 2000 Hz. The impedance change in response to stimulation was recorded over a 15 dB range above threshold and data were fitted with a linear function. Because absolute magnitude of the reflexes varies with individual differences in static
compliance, the ratios of slopes in uncrossed and crossed conditions were calculated within individuals and conditions (right and left ears at 3 frequencies).

Results
Children with suspected APD showed larger slope ratios (uncrossed/crossed), on average exceeding 2. In contrast, slope ratios were smaller in the normal hearing adults and children, on average 1-1.5. There were no significant differences as a function of stimulus frequency or ear.

Conclusion
Larger slope ratios in the children with suspected APD resulted from shallower growth of the amplitude of the crossed reflex with increasing stimulus level when compared to that of the uncrossed reflex. This pattern may suggest reduced neural integrity in the auditory brainstem pathways of many children with APD. The relationship between acoustic reflex abnormalities and functional hearing, especially in noise, may be significant.

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Interaction between cognition and audibility in speech-in-noise hearing in middle aged people
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Background
The launch of UK Biobank (www.ukbiobank.ac.uk) opened a major international opportunity to address an almost limitless range of issues in translational hearing research. Epidemiological data from the 1980s showed that 21% of the UK population aged 40-69 years had a hearing loss (Davis, Hearing in Adults, 1995), with a higher incidence predicted as the population ages. However, recent US evidence has found that as people live longer they retain sensitive hearing later in life (Zhan et al, Am J Epidemiol, 2010, 171, 260-6). No recent comprehensive UK data on the prevalence of hearing disability are available. There has never been a large scale, quasi-random sampling of performance on a speech-in-noise (SiN) test. By comparing self-reported hearing disability in the older (Davis, 1995) and current (Biobank) samples with objective measures of audiometry (older sample), SiN and cognitive performance (Biobank), we will test the hypothesis that a decline in cognition with age will be associated with a decline in SiN hearing that is greater than predicted by audiometry.

Methods
About 500,000 people from across England, Scotland and Wales aged between 40-69 years in 2006-2010 participated in Biobank. About 170,000 completed a specially developed version of the digit triplet test (DTT), an adaptive, high redundancy speech-in-(speech-shaped) noise (SiN) test. In addition to the hearing tests, participants provided data on a wide range of other markers, including age, sex, lifestyle, cognition, employment, socioeconomic, nutrition, medication and other medical variables that have all been linked to hearing. Participants also provided biosamples (blood, saliva, urine) that can be used for testing relationships between hearing and its underlying biology (e.g. genomics and metabolomics). Cognitive tests were fluid intelligence, executive function, processing speed, and auditory (digit recall) working memory.

Results
25.6% reported in a touchscreen questionnaire that they have difficulty with their hearing and 37.5% found it difficult to follow a conversation if there was background noise. Other summary data (www.biobank.ctsu.ox.ac.uk) show mean DTT response thresholds (~6.6 dB S/N) and cognitive test scores in line with expectations. Linking data for Biobank that will enable the hypothesis to be addressed are being released during October 2012.

Conclusion
In addition to the above data, we will present full UK Biobank data on the DTT that will show the detailed relationship between audibility, SiN perception and cognitive performance.

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Test Performance of a Computer Version of the Digits-in-Noise Test for Adult Hearing Screening
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Science University, 4Communication Disorders Technology, Inc., Bloomington, IN, 5Dept. of Speech and Hearing Sciences, Indiana University

Background
The telephone Digits-in-Noise (DIN) hearing screening test has been used in the Netherlands and other European countries since about 2005. Watson et al. (2012) developed an English version of the test with a Middle American dialect. The test uses spoken three-digit sequences presented in speech-spectrum background noise over the telephone. The signal-to-noise ratio for 50%-correct performance is obtained using an adaptive tracking procedure. The purpose of the present study was to evaluate the DIN test for use on a computer using a sound card and headphones. For the purposes of this study, the American English version of the telephone DIN test was modified for use with a computer interface by Watson and Kidd. We were interested in evaluating the use of the computer version of the DIN as a screening test for hearing loss in adults. It was expected that performance might differ somewhat from the telephone version of the test given the greater signal bandwidth available with the headphones used (Sennheiser HDA 200) compared to the telephone, and the new set of recorded sequences selected for this version.

Methods
Forty subjects were recruited for the study; 16 females and 24 males with mean age of 52.2 years. There were 32 ears with normal hearing and 48 ears with sensorineural hearing loss defined as having one or more pure-tone thresholds at 10 frequencies from 250 to 8000 Hz ≥30 dB HL. No subject had an audiometric air-bone gap >10 dB. Following pure-tone audiometric testing in a sound-treated booth, subjects were seated in a quiet room for computerized DIN testing. Subjects followed on-screen instructions for completing a few training trials and then started the test. The test was completed on the right ear and then the left.

Results
For a hearing loss criterion of 3-frequency pure-tone average (1, 2 and 4 kHz) ≥25 dB, the area under the ROC curve (AROC) was 0.95. When a criterion of hearing loss was set at two or more frequencies from 250 to 8000 Hz ≥30 dB HL, the AROC was 0.96.

Conclusion
The computer version of the DIN demonstrated excellent properties for our sample of 32 ears with normal hearing and 48 ears with sensorineural hearing loss. The excellent AROC results (≥0.950) suggest that this version of the DIN should be very useful for adult hearing screening.

Which Children Should be Considered Candidates for Bilateral Cochlear Implantation?
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Background
We are conducting a longitudinal comparison of outcomes for children with bilateral cochlear implants and children with bilateral acoustic hearing aids. The goal is to define a criterion of candidacy for pediatric bilateral cochlear implantation.

Methods
Twenty-seven of the participants use bilateral cochlear implants and have severe-to-profound hearing impairment. Forty-one of the participants use bilateral acoustic hearing aids and have hearing impairment ranging from moderate to profound. Children are assessed twice with an interval of a year between assessments. To date, all children have completed the first assessment, aged 4.3 years on average (range 3.0 to 6.3 years). Participants completed assessments of: 1) language skill; 2) sound-source localization; 3) speech perception in quiet, in random noise, and in babble.

Results
For each outcome measure, a regression function was used to characterize the relationship between unaided hearing level and performance for children with acoustic hearing aids. The distribution of scores for implanted children was used to calculate, for a newly-diagnosed child with a known hearing level, the odds of better performance with implants than with hearing aids. A criterion of candidacy can be defined as the most advantageous hearing level for which the odds of better performance with implants exceed an acceptable ratio, such as 4 to 1.

As predicted, the results varied with the outcome measure. Based on the results of the first assessment, the four-frequency pure-tone average hearing level (PTA) associated with odds of 4 to 1 was 90 dB (speech perception in quiet) and 95 dB (language skills). The remaining outcome measures were not suitable for the proposed method of data
analysis, because there was insufficient overlap of the ranges of scores shown by children with cochlear implants and children with hearing aids.

Conclusion
Based on the results of the first assessment, children with PTA greater than 90 dB should be considered candidates for bilateral cochlear implantation. We will present results from the second assessment at the conference. Supported by Action on Hearing Loss.

Speech Discrimination and Spatial Release from Masking in Toddlers with Cochlear Implants
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Background
Bilateral cochlear implants (BiCIs) have been shown to promote the development of spatial hearing skills in children and adults, but little is known about the role of bilateral stimulation in enhancing language and speech reception in infants and toddlers. In addition, the difference in performance between toddlers with unilateral cochlear implants (UCIs) and BiCIs remains unexplored. We tested two hypotheses:
1. Toddlers with BiCIs will discriminate minimal pair contrasts that differ in voice and/or place of articulation with higher accuracy than toddlers with UCIs because they receive “two looks” at the speech signal.
2. Toddlers with BiCIs will show more spatial release from masking than toddlers with UCIs.

Methods
Toddlers with UCIs and BiCIs (24-36 mo.) were tested. Experiment 1: In a novel Reaching for Sound paradigm, Toddlers were trained to reach for an object whose location the correct stimulus, and were reinforced for correct responses (toy, snack, sticker, etc.). The stimulus consists of the carrier phrase “I’m hiding under” followed by one of the three test words (bee, pea, or key). The locations for the discrimination set are randomized within each group of toddlers (e.g., pea-Left vs. bee-Right), and percent correct was recorded. Experiment 2: Using a computerized 4 alternative forced choice (AFC) task, we adaptively measured speech reception thresholds (SRTs) in quiet and in the presence of a masker whose location was varied between 0° and 90°.

Results
Preliminary results indicate that in Experiment 1 NH toddlers have high %correct scores, i.e., clear discrimination of all speech contrasts, while toddlers with BiCI or UCI have low %correct scores. As hypothesized, toddlers with CIs discriminate voicing contrasts with higher accuracy than place contrasts, potentially due to temporal information being presented more reliably than spectral cues by CI processors. In Experiment 2 NH toddlers demonstrate a benefit when target and masker are spatially separated while toddlers with BiCIs and UCIs are highly variable in their performance on this task.

Conclusion
Speech discrimination results to date suggest that: (1) In implanted toddlers, speech discrimination abilities develop faster for temporally-based speech cues than spectrally-based cues, but thus far there is no measurable difference between UCI and BiCI users. (2) The rate of development in NH toddlers is unknown, because they perform equally well with all stimuli. NH toddlers show a benefit when target and masker are spatially separated while toddlers with BiCIs and UCIs have not.

Acoustic Evaluation of Ambient and Performance Noise in Georgia Aquarium
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Background
Many aquaria hold marine mammal shows for entertainment and educational purposes. Most of these performances are delivered over public announcement (PA) systems. The hearing safety of exhibits and audience is of utmost importance before, during and after the performances. Georgia Aquarium invited the FETCHLAB to ensure that their new dolphinarium met the regulated noise standards. FETCHLAB stands for Facility for Education and Testing of Canine Hearing & Laboratory for Animal Bioacoustics. In addition to testing canine hearing, the FETCHLAB performs noise assessments of animal housing facilities such as kennels and aquaria. We performed both underwater and in-air noise
assessments of the dolphinarium; though the focus of this paper is the in-air acoustic evaluation as it relates to human hearing.

Methods
Sound intensity measurements were made at various locations throughout the dolphinarium theatre and stage. The places to be measured were selected in reference to where patrons of the show would be seated during a performance in the theatre. Sound intensity measures were taken using two calibrated sound level meters (SLMs) - Quest Diagnostics and B&K. Burst measurements (e.g. water effects) were collected using a ‘fast’ SLM response; and measurements of songs and theatrics were collected using a ‘slow’ response. Two sets of recordings were made – ambient and performance noise levels – at 15 locations in the theatre (Figure 1). In addition, noise measurements were performed in the pump rooms to ensure hearing safety of workers and performers.

Results
The recordings revealed average ambient noise levels of 61.4dBA (range: 53-69dBA) and performance noise levels of 96.9dBA (range: 92.6-101.1dBA) for the story track and 98.1dBA (range: 92.3-104.7dBA) for the music track. Noise level in the main pump room was measured at 87.4dBA (range: 86.9-88dBA) and that in the supporting equipment room at 74.1dBA (range: 72.2-76.1dBA).

Conclusion
According to Occupational Safety and Health Administration (OSHA) guidelines, these levels, although high, are well within the safe noise exposure levels as long as the exposure time is limited to 8 hours/day. Use of hearing protective devices (e.g. ear muffs, ear plugs) is recommended for hearing conservation. Annual hearing screenings of workers and annual noise measurements at the dolphinarium are recommended. Future research may focus on the use of noise dosimeters worn by workers to measure the noise dose that they are exposed to in a typical work-day.
Reliability of the Reference Sound Pressure Level (RSPL) for Korean Speech Audiometry
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Background
The reference sound pressure level (RSPL) for speech audiometry is established based on the speech recognition threshold (SRT) measured by standardized words and procedures (ISO 8253-3, IEC 60645-2). The purpose of this study was to examine the reliability of the RSPL for Korean speech audiometry using 36 bisyllabic words which were adopted as the Korean standard words for speech audiometry (KS I ISO 8253-3-2009).

Methods
Subjects were 230 normal hearing persons from 5 universities over the country aged between 18 years and 25 years. Criteria of subjects were type A tympanometry and pure tone thresholds less than 10 dB HL from 250 Hz to 8000 Hz. SRTs were measured by the descending method described in ISO 8253-3 only at the better ear based on the PTA or right ear if PTAs were equal using 36 bisyllabic words which consist of 3 sets and each set has 12 test items. Retest of SRT was performed for the same subject about an hour after the first test. was based on mean difference, correlation, standard error of measurement (SEM) and 95 % confidence interval (CI) were calculated for test-retest reliability of the RSPL for Korean speech audiometry.

Results
Mean SRTs for the test and retest were 23.45 and 23.18 dB SPL with the standard deviations of 3.79 dB and 4.02 dB, respectively, which was not statistically significant (paired t-test, p > .05). Correlation between two tests was significant \( r=0.73, p=.000 \) and SEM was 1.97 dB. Finally, CI of 95% was between 19.51 and 27.39 dB SPL for the RSPL of Korean speech audiometry.

Conclusion
The reliability of the RSPL for Korean speech audiometry is considerably high based on the mean difference, correlation and CI. Therefore, it is suggested that Korean RSPL be 23.45 dB SPL for speech audiometry, which is 3.45 dB greater than English RSPL.
Measuring Cochlear Ischemia using a Combined Dual-Wavelength Laser Speckle Contrast Imaging and Doppler Optical Microangiography System

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\textsuperscript{1}University of Washington, \textsuperscript{2}OHSU

Background
Several hearing disorders such as noise-induced hearing loss, age related hearing loss, sudden sensorineural hearing loss, tinnitus and Ménière’s disease, have been related to cochlear ischemia. Cochlear ischemia is an understudied area of research due to the lack of tools available. It has been difficult to obtain information regarding cochlear ischemia in humans given that the invasiveness of the existing techniques carries the risk of increasing the functional loss. Understanding the mechanisms underlying the pathophysiology of the cochlear microcirculation and oxygenation in a non-invasive manner and in vivo is of fundamental clinical importance, and may enable more effective understanding and management of these disorders.

Methods
A synchronized dual-wavelength laser speckle contrast imaging (DWLSCI) system and a Doppler optical microangiography (DOMAG) system was developed, and used to determine several ischemic parameters in the cochlea due to a systemic hypoxic challenge. DWLSCI can obtain two-dimensional data (the system averages the signal within a small volume of tissue), and was used to determine the changes in cochlear blood flow, and the concentrations of oxyhemoglobin (HbO), deoxyhemoglobin (Hb) and total hemoglobin (HbT) in mice. DOMAG can obtain three-dimensional data, and was used to determine the changes in cochlear blood flow with single vessel resolution.

Results
The changes in the concentrations of Hb, HbO and HbT, as well as the changes in cochlear blood flow were monitored through time. During the hypoxic challenge there was an increase and decrease in Hb and HbO, respectively, and a slight decrease in HbT. The rate of change in the concentrations of Hb and HbO were quantified during and after the hypoxic challenge. The cochlear blood flow decreased during the challenge.

Conclusion
The ability to simultaneously measure these ischemic parameters with high spatio-temporal resolution will allow the detailed quantitative analysis of several hearing disorders, and will be useful for diagnosing and developing treatments.
Miami Otogenetic Program: Implementing genomic medicine in clinical care of deaf patients
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Background
The identification of numerous genes causing NSHL (nonsyndromic hearing loss) along with recent technological advances in “target-enrichment” methods and next generation sequencing (NGS) is now making possible molecular epidemiological studies of genetic deafness. The translation of this knowledge to patient care is, however, lagging. There is an urgent need to bring comprehensive genomic information of individual patients into the “real world” clinical environment. The Miami Otogenetic Program is providing a unique platform to translate laboratory studies to deaf patient care.

Methods
We have collected a unique cohort of multiplex families derived from three sources, the USA, China, and Turkey, suitable for determination of molecular epidemiology of hereditary deafness and for new gene identification. Using target-enrichment/NGS, we are determining 1) the overall frequencies of different forms of genetic deafness in each of the three populations and identify ethnic differences in the distribution of genes for HL, 2) identifying new genes for ARSNHL (autosomal recessive nonsyndromic hearing loss) and ADNSHL using traditional and innovative technologies in the collected families, and 3) creating important Genomic Deafness Database (GDD) and Personalized Sequence Profile (PSP) for the clinical care of deaf patients where data is ranked based on its clinical validity and utility. Our interdisciplinary and collaborative team will conduct outcome evaluation of genetic service on deafness patient care in our diversity populations.

Results
We have established the Miami Otogenetic Program which includes both research and the clinical components. The infrastructure of our multidisciplinary otogenetics team will be presented along with our utilization of testing algorithms when evaluating patients with SNHL. We have collected DNA samples from over 800 probands from multiplex families with no mutations in the common deafness genes from three unique cohorts. In 60% of the small multiplex families, we have identified mutations in known deafness genes. The remaining 40% have mutations in other yet-unidentified deafness-causing genes. We have identified the genes for the DFNA41 and DFNB84 loci in large multigenerational families as well as three new deafness genes for NSHL in small multiplex families.

Conclusion
The combined target-enrichment/next generation sequencing (NGS) and WES (whole exome sequencing) is a powerful tool in the identification of new deafness genes in small multiplex families and large multi-generational families. The multidisciplinary team approach is an effective way to bring the sequencing data to clinical practice for the clinical diagnosis and management of deaf and hard-of-hearing families.

Otitis Media Impacts Hundreds of Middle and Inner Ear Genes
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1OHSU

Background
Otitis media is known to alter gene expression of selected cytokine, mucin and ion homeostasis middle ear (ME) and inner ear (IE) genes in the mouse model. However, whole mouse genome studies of the gene expression in a setting of otitis media have not previously been undertaken. Ninety-nine percent of genes in the mouse are shared in the human, so these studies are relevant to the human condition.

Methods
To assess inflammation-driven processes in the mouse ME and IE, gene chip analyses were conducted on mice treated with trans-tympanic heat-killed Hemophilus influenza. ME and IE tissues were harvested at 6 hours, RNA extracted, and 7-8 samples for each treatment processed on the Affymetrix 430 2.0 Gene Chip for expression of its 21,816 genes. This represents approximately 70% of the entire mouse genome.

Results
Results were statistically analyzed for expression of each gene against control (untreated) mice. Significant alteration of gene expression was seen in 2,355 genes, 11% of the genes tested and 8% of the mouse genome. Significant ME and
IE upregulation (fold change >1.5, p<0.05) was seen in 1,081 and 599 genes respectively. Significant ME and IE downregulation (fold change <0.67, p<0.05) was seen in 978 and 287 genes respectively. Twenty-nine percent of IE genes upregulated were unique to the IE, while 60% were unique to the ME. Forty-two percent of the IE genes downregulated were unique to the IE, while 83% were unique to the ME. These studies show hundreds of ME and IE genes are affected by inflammation. While otitis media is widely believed to be an exclusively middle ear process with little impact on the inner ear, the IE gene changes noted in this study were numerous and discrete from the ME responses. This suggests that the IE does indeed respond to otitis media and that the IE response to ME inflammation is a distinctive process from that occurring in the ME. Numerous new genes, previously not studied, are found to be affected by inflammation in the ear.

**Conclusion**

Whole genome analysis via gene chip allows simultaneous examination of expression of hundreds of gene families influenced by inflammation in the ME. Discovery of new gene families affected by inflammation may lead to new approaches to the study and treatment of otitis media.

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**Biofilm structures in Infants with Silent Otitis Media and Tympanogenic Meningitis**

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¹University of Minnesota

**Background**

Silent otitis media is defined as chronic pathologic conditions that are "undetected" behind an intact tympanic membrane. Because pathologies in the middle ear are undetected, there is a lack of clinical treatment increasing the risk of complications. Since the initial description of biofilms in chronic infections, there has been increasing evidence supporting the related role of biofilms on disease progression and persistence of inflammation in chronic otitis media. Our hypothesis is that there is an association between the presence and location of biofilms in the middle and inner ear of patients with silent otitis media and tympanogenic meningitis.

**Methods**

Thirty-one cases with meningitis were screened to select those with tympanogenic meningitis. Inclusion criteria for tympanogenic meningitis were acute meningitis with histopathological evidence of chronic otitis media, and no other source of infection. Pathologic changes of the middle and inner ears were noted. We defined biofilm as bacterial aggregates encapsulated within a surface that were frequently associated with a fibrous matrix and contained inflammatory cells. In order to detect the bacteria in the specimens and their aggregates, gram staining, matrix as well as dead-bacteria staining has been performed.

**Results**

Seventeen temporal bones, from 9 cases (2 females and 7 males), ranging in age from 5 to 23 months, met our criteria of tympanogenic meningitis. Of these, 82% had biofilms that contained bacteria within fibrous matrix as well as the presence of polymorphonuclear leukocytes and mononuclear cells. Biofilm-like structures were located in 1 anatomical region in 1 ear and multiple regions in 16. Most common locations were the areas near the oval and round windows and the epitympanum, facial recess, and supratubal recess. Biofilms within the inner ear were observed in the scala tympani and modiolus in 9 ears. Labyrinthitis was seen in all ears as were intact tympanic membranes.

**Conclusion**

There is an association between the presence of biofilm in the ear and tympanogenic meningitis in infants. Chronic pathologic changes in the middle ear, behind an intact tympanic membrane and the lack of ear symptoms, may result in potentially serious sequelae and complications in infants. The clinician should, therefore, be aware that an intact tympanic membrane does not necessarily preclude the presence of pathologic changes such as granulation tissue, cholesterol granuloma, fibrosis and/or biofilm structures in the middle ear cleft, and that the clinical definition of chronic otitis media may not cover the majority of infants with this condition.
Limitations of hearing screening in newborns with a pendrin mutation
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Background
Pendrin gene mutations are the most common cause of congenital hearing loss in East Asia. Because hearing loss caused by the pendrin mutation tends to have a delayed presentation, universal newborn hearing screening (UNHS) examinations are poorly predictive of long-term hearing ability in these patients. We examined associations between UNHS and long-term hearing outcome in deaf patients with biallelic pendrin mutations.

Methods
Forty-three patients with severe to profound hearing loss were recruited. Patients had an enlarged vestibular aqueduct and biallelic PDS mutations. Seventeen deaf patients without a PDS mutation were recruited as a comparison group. We reviewed children’s hearing loss history using medical records and parent interviews.

Results
Fourteen of 43 children (32.6%) with a pendrin mutation received a UNHS test. Four of these 14 children (28.6%) passed in both ears, while six (42.9%) passed in one ear, and four (28.6%) were referred in both ears. In contrast, 15 of 17 children (88.2%) without a PDS mutation were referred bilaterally. The age at confirmation of bilateral hearing loss in bilateral pass children was 31.5±17.9 months, which was significantly later than the age for bilateral refer children (1.75±0.96 months) (p=0.045).

Conclusion
The UNHS is not an accurate tool for predicting long-term hearing loss in patients with a pendrin mutation. We recommend that genetic screening be combined with UNHS, particularly in communities with a high prevalence of pendrin mutations, to better identify children in need of early habilitation.
Table 1. Clinical characteristics of patients with PDS mutation

<table>
<thead>
<tr>
<th>Patient number</th>
<th>Gender</th>
<th>Age at which hearing loss was confirmed (months)</th>
<th>Pendrin mutation type</th>
<th>Screening device</th>
<th>Type of UNHS</th>
<th>Setting threshold</th>
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<tr>
<td>1</td>
<td>F</td>
<td>54</td>
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<td>VIASYS</td>
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<td>GSI-Audera</td>
<td>ABR</td>
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<td>4</td>
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*The patient 3 performed the ABR at age 18 months and was confirmed with normal at age 24 months. The patient 8 performed the ABR at age 2 months and was confirmed with one side hearing loss at age 16 months.
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Otology in the classical world -- 500 BCE to 200 CE
Robert Ruben
1Albert Einstein College of Medicine

Background
This presentation is part of a history of otology using a uniform subject matter framework across time. Information is accessible through hypertext based upon the subject.

Methods
Original source material is searched using uniform criteria for subject categories.

Results
The Hippocratic corpus (ca 460BCE -- 377BCE) describes conditions interpretable as otitis media which appears as a lethal disorder, without interventions. Deafness is cited among sequelae of other disorders in which there is recovery. Celsus (25BCE -- ca 50CE) emphasizes the seriousness of ear infections, especially for young and old. Serous Otitis Media: Hippocratic writings note conditions interpretable as SOM, associated with southerly winds and/or dampness, causing a fullness of the head and deafness. Acute onset deafness associated with apparent systematic infection is mentioned. These cases do not indicate initial ear infections but the ear is affected in the illness’s course. SOM is not recognized as an entity.
Barotrauma: Aristotle (384 BCE -- 322BCE) appreciates the relationship of middle ear pressure to external pressures, noting that tympanic membrane rupture in deep sea diving is preventable by attaching a sponge to the external ear. He attributes the protection to blocking of sea water, although the sponge’s evident effect is lessening pressure on the tympanic membrane caused by lower pressure in the middle ear cleft.
Deafness: Greeks and Romans provide no etiologies for congenital deafness. Aristotle recognizes hearing’s essential role in spoken language development, and notes greater morbidity of congenital and early onset deafness compared with blindness. Mutism is attributed to inability to use the tongue. Plato (ca 360BCE) describes the use of sign language by the deaf.
Ear Surgery: Celsus describes hearing restoration in congenital or acquired external canal stenosis using caustic and/or incision with stenting. His techniques for repairing pinnae injuries are much like contemporary practice.
Medical providers: The general population received medical care from general physicians, some paid, in part, by their communities. The wealthy had personal physicians. Rome and Alexandria had otologic specialists adept at removing external canal foreign bodies and caring for pinnae injuries.

Conclusion
The seriousness of ear infection was recognized, with no available interventions. Care existed for external canal stenosis, pinnae injuries and external canal foreign bodies. Hearing’s essential role in spoken language acquisition was known, with lack of speech attributed to tongue dysfunction. Sign language of the deaf was recognized.
Ototoxicity of Burow's solution on the guinea pig cochlea
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1Department of Otorinolaryngology, Fukuoka University, school of Medicine, 2Nishi Fukuoka Hospital

Background
The main ingredient of Burow's solution is 13% aluminum acetate and it has been topically used worldwide for the treatment of chronic discharging ears. However, ototoxicity of this solution is not well understood. The aim of the present study is to evaluate the ototoxicity and the bacteriostatic activity of Burow's solution with different concentration.

Methods
Using guinea pigs, compound action potentials (CAPs) of the eighth nerve were measured before and 30 minutes and 24 hours after the application of the Burow's solution in the middle ear cavity. The bacteriostatic activity of Burow's solution was studied using a disk diffusion assay.

Results
Use of the original Burow's solution (pH 3.5) for 30 minutes caused a significant reduction of CAP threshold to click sounds. A 2-fold diluted Burow solution (pH 4.4) for 30 minutes caused no reduction in CAP threshold. Burow's solution, pH adjusted to 4.5, caused no changes in CAP threshold at 30 minutes. At 24 hours, original Burow's solution (pH 3.5) caused complete abolition of CAP. Both the original concentration and the 2-fold diluted Burow's solution showed bacteriostatic activity against all bacteriae.

Conclusion
Burow's solution is ototoxic in the guinea pig when applied in the middle ear cavity for 30 minutes or longer. In the clinical settings, it is advisable to avoid allowing the solution to contact the round window membrane for extended duration. We should be aware of anatomical differences of the round window membrane in rodents and human.

References
Postoperative Outcome of Cochlear Implantation in Children with Cochleovestibular Malformations

Teru Kamogashira¹, Akinori Kashio¹, Erika Ogata¹, Yusuke Akamatsu¹, Shotaro Karino¹, Takashi Sakamoto¹, Akinobu Kakigi¹, Shinichi Iwasaki¹, Tatsuya Yamasoba¹

¹Department of Otolaryngology, Faculty of Medicine, University of Tokyo

Background
Malformations of the inner ear were initially considered to be a contraindication to cochlear implantation (CI). However, most cases with inner ear malformations have demonstrated significant improvements in speech recognition skill after CI. Recently, Sennaroglu proposed a new classification system for inner ear malformations on the basis of radiological findings. In this study, we aimed to assess the outcomes after CI in children with a variety of inner ear malformations classified according to the Sennaroglu's classification.

Methods
Data of children who underwent CI at The University of Tokyo Hospital from May 1999 to April 2012 were collected. The cochleovestibular malformation in each case was evaluated using computed tomography and magnetic resonance imaging. Subjects were divided into 7 groups according to Sennaroglu's classification. The Meaningful Auditory Integration Scale (MAIS) scores and the Meaningful Use of Speech Scale (MUSS) scores were analyzed before and 12 months after CI. The incidence of CSF gusher and facial nerve stimulation as well as the number of inserted electrodes were also compared among groups.

Results
Of the 164 children that received CI, 28 had cochleovestibular malformations that were classified as follows: incomplete partition type I (IP-I) (n = 4); incomplete partition type II (IP-II) (n = 15); enlarged vestibular aqueduct (EVA) (n = 2); and common cavity (CC) (n = 3). Other cases that did not match with Sennaroglu's classification included the following: enlarged internal acoustic meatus (n = 2); narrowed internal acoustic meatus (n = 1); and lateral semicircular canal aplasia (n = 1). Three children with IP-II had other complications, including Waardenburg syndrome, Goldenhar syndrome, and CHARGE syndrome, respectively. The average MAIS scores before and after the CI were 11 and 30 in cases of CC, 27 and 37 in those of EVA, 9 and 23 in those of IP-I, and 11 and 24 in those of IP-II. The corresponding MUSS scores were 10 and 26 in cases of CC, 28 and 34 in those of EVA, 7 and 19 in those of IP-I, and 9 and 21 in those of IP-II. Cerebrospinal gusher occurred in 1 CC case and 5 IP-II cases. All electrodes were successfully inserted in all cases except in 1 IP-II case.

Conclusion
The Sennaroglu's classification does not impact the outcome of MAIS and MUSS. However, the risk of CI complications is high in patients with cochleovestibular malformations.
Computed tomography-based preoperative evaluation of round window niche visualization from facial recess

Akinori Kashio1, Akinobu Kakigi1, Shotaro Karino1, Takashi Sakamoto1, Shinichi Iwasaki1, Tatsuya Yamasoba1
1University of Tokyo

Background
The introduction of new electrode array designs and the increased emphasis on preserving residual hearing for lector-acoustic stimulation has recently focused attention on the use of the round window membrane (RWM) approach or extended-RWM approach for cochlear implant (CI) electrode insertion. The visualization of round window niche (RWN) from the facial recess would be an important factor for evaluating accessibility of RWM at the CI surgery. In this study, we aim to preoperatively evaluate RWN visibility by using high resolution computed tomography (HRCT).

Methods
We retrospectively reviewed data from 46 CI patients who underwent HRCT of the temporal bone and whose surgical video was recorded. The axial slice showing RWN was selected and a line was drawn along the posterior wall of the bony external auditory canal (line A). Another line was drawn parallel to line A that made contact with the vertical segment of the facial nerve (line B). The band within line A and B was extended toward the RWN and the relation between the band and the RWN was classified into 3 types: [1] almost no contact with the band, [2] fully included in the band, and [3] partially included in the band. In this procedure, we eliminated 3 cases owing to tortuous external auditory canals. Next, the visibility of the RWN was evaluated and classified into 3 types: (A) almost impossible to visualize, (B) fully visible, and (C) Partially visible.

Results
Ten cases were classified as [1] by CT and all of them were classified as (A) by the surgery video record; 14 out of 16 cases classified as [2] by CT were classified as (B) by the surgery video record. The rest of 2 cases were classified as (C); 16 out of 17 cases classified as [3] by CT were classified as (C) by the surgery video record and the remaining 1 case was classified as (A).

Conclusion
RWN visibility and the preoperative HRCT findings have showed high correlation. Our method for the preoperative evaluation of the RWN has the potential to become a useful tool for CI surgery.

EUSTACHIAN CUSHION FEATURES DO NOT CORRELATE WITH OTITIS MEDIA

N Wendell Todd1, Mina Tran1
1Emory

Background
Diminutive inferiorly positioned Eustachian cushions, as viewed with fiberoptic nasopharyngoscopy, have been associated with otitis media. Our objectives are to describe Eustachian cushion position and features in clinically normal cadaver specimens, and address the hypotheses that bilateral symmetry exists and that Eustachian cushion position and features are unrelated to the small mastoid size indicator of otitis media in childhood.

Methods
From 41 adult cranial specimens without clinical otitis, rectilinear digital photographs were made of the lateral nasopharyngeal walls, both en face directly laterally and viewing 45 degrees posterior-lateral. Features were categorized, and distances measured. Mastoid sizes were quantified on radiographs.

Results
Though good repeatability of assessments and bilateral symmetry of Eustachian cushion features and positions were found, there was no relation of any of the studied Eustachian cushion parameters with mastoid size.

Conclusion
No difference in Eustachian cushion architecture was found to correlate, in these clinically normal crania, with the small mastoid size indicator of childhood otitis media.
Ear-Canal Reflectance as a Screening Tool for the Diagnosis of Superior Semicircular Canal Dehiscence

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1Speech and Hearing Bioscience and Technology, Harvard-MIT Division of Health Sciences and Technology, 2Eaton-Peabody Laboratory, Massachusetts Eye and Ear Infirmary, Boston, MA, USA, 3Department of Otology and Laryngology, Harvard Medical School, Boston, MA, USA, 4Department of Otorhinolaryngology - Head and Neck Surgery, University Medical Center, Utrecht, 5Department of Radiology, Massachusetts Eye and Ear Infirmary, Boston, MA, USA

Background
Ear-canal-reflectance (ECR) was measured in patients with superior semicircular canal dehiscence (SCD) to determine whether ECR coupled with an appropriate detection algorithm can be used as a non-invasive, fast, and easy screening test for SCD. The aim of such screening is to determine if the diagnosis of SCD (and other third-window lesions) should be considered for a patient with vestibular and/or auditory symptom(s) and whether a further workup (such as computed tomographic (CT) imaging and balance testing) should be pursued.

Methods
We measured ECR in 32 ears diagnosed with SCD by high-resolution CT imaging, utilizing specialized methods for measuring size and location of the dehiscence, and compared these measurements to ECR measurements made in normal ears. An analysis algorithm was introduced that detects possible SCD using unique features of ECR measurements that are common in ears with SCD and less common in normal ears. We: 1) evaluated sensitivity and specificity of ECR and our algorithm in the diagnosis of patients with SCD regardless of symptomatology (i.e. with or without hearing loss, dizziness, or abnormal auditory sensations), 2) compared ECR to another non-invasive mechanical measurement – umbo velocity measurements by laser Doppler vibrometry – which has been shown to aid in differentiating various pathologies associated with conductive hearing loss (Rosowski et al. 2008, Nakajima et al. 2011) and 3) correlated specific parameters of ECR measurements to SCD size and location and the air-bone gap seen on the audiogram.

Results
SCD was commonly associated with a notch in ECR near 1 kHz. We assessed the ability of our algorithm to detect SCD based on this ECR notch; various detection thresholds yielded sensitivities of 0.926–0.852, specificities of 0.690–0.724, negative predictive value of 0.925–0.913 and positive predictive value of 0.581–0.589. There was no similar stereotypic feature visible in umbo velocities measured in ears with SCD cases. Statistical analysis showed that there were significant correlations between air-bone gap and ECR notch size, as well as between air-bone gap and an increase in low-frequency umbo velocity magnitude, but no correlations were found between notch size and the size and/or location of the dehiscence.

Conclusion
Our findings suggest that notches in ECR measurements can be used to screen patients for third-window lesions such as SCD in the early stages of a diagnostic workup.

Development of head and neck cancer xenograft animal model for in vivo anti-cancer drug screening

In Seok Moon1, Mee Hyun Song2, Jae Young Choi1

1Yonsei University College of Medicine, 2Kwandong University College of Medicine

Background
The development of a relatively simple, reliant and cost-effective animal test will greatly facilitate drug development. In this study, our goal was the establishment of a rapid, simple, reproducible, and live zebrafish head and neck squamous cancer cell (HNSC) and thyroid cancer cell xenograft model for anti-cancer drug screening.

Methods
We optimized the conditions for the cancer cell xenograft in terms of injected cell numbers, incubation temperature and time. We used cell lines as follows: HNSC (Cal27, FaDu) & Tyroid cancer (FTC33, TPC1). Cancer cells were stained with a mCherry fluorescent protein prior to injection into the 30hpf zebrafish larvae. Cancer cells were counted by microscopy, suspended in 10% FBS and 200 cells were injected into the center of yol sac using a micro-injecting system. Injected embryos were transferred to E3 media and subsequent tumor formation and cancer cell dissemination was observed under fluorescent microscopy for 5 days. To test our new model, we carried out further validation. Xenografted embryos
were transferred in a 24 well plate (3 embryos/well) and treated with several known anti-cancer drugs (RAD001 and others).

Results
MFP-tagged cancer cells were successfully grafted into the yolk sac of zebrafish embryo without immunosuppressant treatment. 2 days after injection, xenografted cell formatted tumor mass in all cell lines. 4 days after injection, dissemination from the yolk sac to the tail can be observed from in FaDu line under fluorescent microscope. Differences in injected cell numbers were reflected in the speed of tumor formation and rate of dissemination from the xenograft site. The preliminary test was performed using 1 μM of RAD001 and others. RAD001 produced > 20% reduction in the cancer cell number of embryos.

Conclusion
Our in vivo animal model system should greatly facilitate drug development for cancer therapy because of its speed, simplicity, reproducibility and live imaging.

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p75NTR-mediated apoptosis in vestibular schwannoma (VS) cells depends on subcellular localization of the intracellular domain
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1University of Iowa Department of Otolaryngology

Background
Our work focuses on p75NTR signaling in neoplastic and non-neoplastic Schwann cells (SC). Post-activation cleavage of p75NTR yields an intracellular domain (ICD) in the cytoplasm that induces apoptosis in non-neoplastic SCs. We recently showed that the ICD of p75NTR localizes to the nucleus in auditory SCs that have re-entered the cell cycle following deafness. Further, work in PC12 cells demonstrated that nuclear-localized p75NTR-ICD regulates cyclin E1 expression. These observations suggest that subcellular localization of p75NTR-ICD regulates SC cell cycle entry and survival.

Methods
p75NTR expression in VS tissue, relative to normal nerve, was quantified by qRT-PCR and western blot. To explore the contribution of p75NTR signaling in VS cell proliferation and death, we generated adenoviral vectors that express full-length (FL) p75NTR, p75NTR-ICD, p75NTR-ICD targeted to the cytoplasm with a nuclear export signal (NES) peptide, or p75NTR-ICD targeted to the nucleus with a nuclear localization signal (NLS) peptide. Primary human VS cultures derived from acutely resected tumors were transduced with these viral vectors or treated with proNGF, a high affinity p75NTR ligand. TUNEL staining and caspase-3 cleavage were used to demonstrate apoptosis. Proliferation was determined by EdU uptake. Finally, transcriptome analysis of VS cultures treated with proNGF was performed using Nimblegen microarrays.

Results
Compared to normal nerve, VSs express a ~3-fold increase in p75NTR. Based on immunostaining, the ICD of p75NTR localizes to the nucleus in EdU-positive VS cells in vitro as in non-neoplastic SCs. Treatment of VS cultures with proNGF resulted in an anti-apoptotic response in VS cells while non-neoplastic SCs responded with increased apoptosis. Expression of FL p75NTR and NES-p75NTR-ICD both increased the percent of TUNEL-positive VS cells compared with cultures transduced with an empty vector. By contrast, expression of NLS-p75NTR-ICD failed to induce VS cell apoptosis. Treatment of VS cultures with proNGF or transduction with the various p75NTR isoforms did not alter the percent of EdU-positive VS cells. Microarray data are currently being analyzed.

Conclusion
VS cells express high levels of p75NTR yet are resistant to proNGF-mediated apoptosis. The ability of p75NTR to induce VS apoptosis appears to depend on the subcellular localization of the ICD. Activation of p75NTR with proNGF altered expression levels of several genes potentially related to neoplasia in VS cells.
Endolymphatic Hydrops in Histopathologic Otosclerosis and Surgery Effect?
Christina Forshell Hederstierna\textsuperscript{1}, Sebahattin Cureoglu\textsuperscript{2}, Vladimir Tsuprun\textsuperscript{2}, Michael Paparella\textsuperscript{2}
\textsuperscript{1}Karolinska University Hospital, \textsuperscript{2}University of Minnesota

Background
Otosclerosis is a disease of the bony labyrinth and stapes, causing a remodeling of the bone structure. In clinical cases, the affected motility of the stapes causes a conductive hearing loss, in one or both ears. However, patients with otosclerosis often experience other symptoms that are not typical of the disease, e.g. symptoms of tinnitus, fullness of the ear, vertigo and dizziness. These symptoms are more often characteristics of Meniere’s disease and related conditions, where the basis for the symptoms is considered to be endolymphatic hydrops.

Methods
We studied the complete collection of temporal bones with histological otosclerosis at the University of Minnesota Otopathology laboratory. 188 ears from 104 subjects with histologic otosclerosis were examined for the degree of cochlear and saccular hydrops and correlated with presence or absence of surgical intervention.

Results
In 4 of these cases only one ear was available for examination. 84 of the remaining 100 cases were bilateral and 16 were unilateral; 6 in the left and 10 in the right ear. In total, 167 ears with otosclerosis had quantifiable cochlear hydrops and 150 ears had quantifiable saccular hydrops according to our specified rationales for grading.

Conclusion
We found that histological otosclerosis is associated with a notable prevalence of both cochlear and saccular hydrops. The number of otosclerotic foci correlates strongly with the degree of cochlear and/or saccular hydrops; the prevalence of both types of hydrops appears lower in ears that have had otosclerotic surgery.
### Contribution of the incudo-malleolar joint to middle-ear sound transmission

**Rahel Gerig**¹, Jae Hoon Sim¹, Christof Röösli¹, You-zhou Xie¹, Sebastian Ihrle², Alexander M. Huber¹

¹University Hospital Zurich, ²University of Stuttgart

**Background**

The incudo-malleolar joint (IMJ) in human middle ears is considered to be the diarthrodial joint binding the malleus and the incus. Investigations on measurement of IMJ mobility have been conducted during the last several decades, and the IMJ in human is generally accepted to be mobile, especially at high frequencies. However, the behavior of the IMJ has not been fully characterized, and its roles in sound transmission are still questionable. Furthermore, the supporting band apparatus of the IMJ may have a relation to age-related hearing loss, but this has not been demonstrated. Characterizing IMJ motions systematically and developing a comprehensive mechanical model of the IMJ can answer questions about its anatomical features and function, and serve as a base for the development and optimization of middle-ear implants.

**Methods**

Spatial motion of the middle-ear ossicles in fresh temporal bones (TB) was measured using a scanning laser Doppler interferometry (LDI) system. Three-dimensional (3-D) motion components of each middle-ear bone were calculated from the measurements, using a custom-made algorithm. The measurement and calculation were repeated with the IMJ experimentally fixed, in order to quantify the transmission losses caused by the immobility of this joint. The spatial motion components were registered into an anatomical frame, which was obtained from micro-CT images. Four pieces of reference wires were attached to the TB close to the measurement site, and their outlines were used as references to obtain the relation between the LDI measurement frame and the anatomical frame.

**Results**

While the malleus and the incus moved as one rigid body with the fixed IMJ, relative motion between the two middle-ear bones was observed with the non-fixed IMJ, especially at high frequencies above 2 kHz. The pattern of the stapes spatial motion also showed differences between the non-fixed and fixed conditions of the IMJ.

**Conclusion**

The spatial modes of the middle-ear ossicles are affected by the IMJ condition, especially at high frequencies. Therefore, the IMJ is presumed to contribute to the middle-ear sound transmission and thus air-conductive (AC) hearing.

### Alternative Lever Arms for Mobile and Fused Ossicular Chains

**Ryan Jackson**¹, Allison Zemek¹, Hongxue Cai¹, Charles Steele¹, Sunil Puria¹

¹Stanford University

**Background**

Several publications have argued that morphometric properties of the ossicular chain play an important role in determining force gains, inertial impedance, and, consequently, the frequency range of hearing in mammals. More recently, Puria and Steele (2010) proposed that at high frequencies the vibration of the ossicles should be predominantly rotational about their individual principal moment-of-inertia axes, rather than their anatomical axis, which is defined by the malleal anterior and incudal posterior ligaments. Such vibrations could significantly reduce the mechanical impedance of the ossicular chain and enable efficient sound transmission to the cochlea. Here the first, second, and third principal axes (PA1, PA2, and PA3 respectively) were compared to the anatomical axis (ANA) and another axis of interest, the malleus-incus complex center-of-mass axis (COMA). Lever arms were calculated for each axis to estimate primary motions that should be observed experimentally if rotation occurs about each of these axes, provided the ossicles behave primarily as rigid bodies. Further, the ratio of the malleus-lever-arm to incus-lever-arm was calculated to estimate middle ear gain as an ideal transformer ratio.

**Methods**

Cadaveric temporal bones from three species with mobile malleus-incus joints (MMIJ): gerbil (n=6), cat (n=2), and human (n=1), and two species with fused malleus-incus joints (FMIJ): guinea pig (n=6), chinchilla (n=2) were scanned and segmented to analyze ossicular chain morphology. Analysis of lever arms, moments-of-inertia, and tangential linear velocity directions for each animal was performed.

**Results**

For species with MMJ, the average values of malleus-to-incus lever arm ratios were increased when the incus rotated about its PA1 for any malleus axis of rotation (increases of 5x for gerbil, 2.5x for cat, and 3.5x for human). A similar maximization was found for species with FMIJ, when the malleus and incus rotated about their combined PA1 (increases...
of 2x for guinea pig and 1.3x for chinchilla). It was also found that for either MMIJ or FMUJ, rotation of the incus about PA1 imparts force to the stapes approximately in the piston direction.

Conclusion
Examination of the lever arm ratios indicated that in the mammals studied, rotation of the incus about its PA1 is significantly greater than is possible with rotation strictly about ANA (or COMA). This could indicate a novel mechanism for increased middle ear gain.

[Work supported in part by NIDCD/NIH Grant R01 DC005960.]

Estimation of the orthotropic elastic properties of the eardrum using a pressurization technique
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¹Western University, ²London Health Sciences Centre

Background
Previously, we reported a method based on indentation testing and optimization of a finite-element (FE) model to estimate the orthotropic elastic parameters of the pars tensa in situ. This approach is challenging to use because of the curved shape of the pars tensa which requires special care during experimentation to keep the indenter perpendicular to the local surface at the point of contact. In addition, FE simulations involve complicated contact modeling. A new method is presented here in which pressurization is used instead of indentation.

Methods
Measurements were made on three rat eardrums with immobilized ossicular chains. A pressurization system was used to apply quasi-static pressures up to 4 kPa to each eardrum. The resting and deformed shapes of each eardrum were measured using a Fourier transform profilometer, a non-contacting optical device for shape measurements. To simulate the pressurization experiment, an FE model was constructed for each eardrum from the resting shape data. The orthotropic parameters of each model pars tensa were optimized so that simulated deformed shapes matched measured ones. A variant of the Nelder-Mead simplex method that is formulated to handle constrained multivariable optimization was used, and the optimal values were taken to be estimates for the actual orthotropic parameters.

Results
The estimated parameter values are circumferential elasticity of 37.9 ± 0.12 MPa, radial elasticity of 48.0 ± 0.01 MPa and in-plane shear modulus of 22.0 ± 0.01 MPa. The average estimated values for the radial and circumferential elasticities are within the range reported in the published literature on tensile testing of tissue strips cut from the pars tensa; however, tensile testing cannot be used to estimate the in-plane shear modulus.

Conclusion
The new pressurization-based approach is simpler to use than the indentation-based method both in terms of measurement and modeling, and yields estimates for circumferential and radial elasticities that are comparable to those published by other groups. Moreover, the results are highly repeatable as indicated by the low standard deviations.
Characterization of Ear Canal Sound Field in Forward and Reverse Transmission
Jeffrey Cheng¹, Michael Ravicz¹, Cosme Furlong¹,², John Rosowski¹,³
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Background
Sound propagation from the external environment through the human ear canal (EC) to the tympanic membrane (TM) can be described fairly well in terms of plane wave propagation at least at frequencies below 15 kHz (Stinson 1985). However, the sound field generated inside the EC by motion of the TM in reverse transmission (for instance, by otoacoustic emissions) has not been well studied. We have reported preliminary measurements showing that (1) the relationship between TM displacement and ear canal pressure differs in both magnitude and phase between forward and reverse transmission (Cheng et al., 2011), and (2) there is little correspondence between the nonuniform TM surface motion in forward transmission and the nearly uniform pressure distribution inside the EC below 15 kHz (Cheng et al., 2012).

Methods
In this study, we compare the distribution of sound pressure inside the EC for both forward and reverse sound transmission. For forward transmission, sound was delivered from an earphone to the opening of an artificial EC (with dimensions similar to the human ear) that was coupled to the tympanic ring of a temporal bone at a 45-degree angle to mimic sound propagation inside the actual human EC; for reverse transmission, the incus body near the long process was stimulated by a piezoelectric actuator (the EC entrance was open). Sound pressure was measured in a 1-mm grid in a plane parallel to and within 1 mm of the tympanic ring with a probe tube microphone. Sound pressure was also measured in a set of planes normal to the EC axis but more distant from the TM ring.

Results
In forward sound transmission the sound pressure was approximately uniform over both the EC normal and the tympanic ring planes at frequencies up to 15 kHz. In reverse transmission the sound pressure varied within the tympanic ring plane at frequencies as low as 1 kHz, but was uniform throughout the EC normal planes. The pressure variations along the tympanic ring plane in reverse transmission tend to be frequency dependent, and the pressure from reverse transmission is significant lower than those from the forward case with the open EC.

Conclusion
Our results suggest the presence of evanescent waves from TM motions in reverse transmission which may influence assessment of oto-acoustic emissions measurements in real ear canals.

Imaging the Human Tympanic Membrane Motion Using Scanning Laser Vibrometry and Finite Element Model
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Background
Sound-induced tympanic membrane (TM) vibration has been studied through the motion of the umbo which was measured by laser Doppler vibrometry or analyzed using finite element (FE) models for decades. Recently, some preliminary studies were conducted to measure the full-field TM surface motion using stroboscopic holographic interferometry and optical coherence tomography. The technology of imaging human TM motion has the potential to be a clinical diagnosis tool for middle ear diseases. However, how is the TM surface motion related with the middle ear transfer function in normal and disordered ears is still not clear.

Methods
In this study, a scanning laser vibrometer (PSV-400, Polytec) was used to measure and image the full-field surface motion of the TM sample from six fresh human temporal bones. Both displacement amplitude and phase were measured under the normal and disordered conditions with different middle ear pressure or fluid levels across 200 to 8,000 Hz. A completed FE model of the human ear, including the ear canal, middle ear and spiral cochlea, which was published recently by our group (Zhang and Gan, 2011) was used to simulate the measurement and derive the holographic images of TM motion.

Results
The mode shape and displacement phase of the TM at low (below 1,000 Hz), medium (between 1,000 and 4,000 Hz) and high frequency ranges (above 4,000 Hz) under normal and disordered conditions were acquired from the temporal bone.
experiments and predicted by the FE model. The complex mode shape and standing wave were observed on TM at high frequencies. The relationship between the mode shape of the TM and the middle ear transfer function was analyzed.

Conclusion
The scanning laser vibrometry is capable to measure and image the full-field motion of the human TM. The holographic images of TM can be clearly distinguished between the normal ear and the middle ear with effusion at high frequencies. The data reported in this study may help better understand the complex TM motion in relation to its acoustic transfer function. (Work supported by NIH R01DC011585)

The Effect of Static Force during Stimulation of the Round Window with the Floating Mass Transducer
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Background
The round window (RW) stimulation has developed to a standard procedure in the treatment of sensor neural hearing losses in combination with middle ear pathologies since Colletti et al. successfully implanted the first series of patients in 2005. Despite encouraging clinical results there is still a high variability in hearing improvement results. Two key requirements seem to play an important role for obtained sufficient sound transmission. First the FMT must be centered onto the round window membrane and secondly the back of the FMT must be supported elastically in the hypotympanic area to apply the necessary contact pressure.

The goal of our study was to investigate the influence of a controlled axial static force on transmission efficiency and variability

Methods
We investigated experimentally the effect of the Round Window Coupler (RWC) and the interposition of an artificial material (TUTOPATCH®) in fresh human temporal bones at constant contact force of ~2mN. The equivalent sound pressure output was determined by comparison of sound evoked to RW stimulation evoked stapes footplate displacement. Results were compared to direct attachment of the FMT to the RW membrane without the interposition.

Results
The obtained mean equivalent sound pressure for all four conditions was not significantly altered at frequencies ≥ 3 kHz. Compared to the direct FMT stimulation the coupler in combination with TUTOPATCH® lead to a statistically insignificant decrease of approx. 5 dB, whereas the use of TUTOPATCH® or the coupler alone resulted in smaller changes. At low frequencies (≤ 2 kHz) the effect of the coupler in combination with the interposition material was more pronounced, leading to an average increase in output of 12.2 dB (500 Hz – 2kHz). The improvement in the output was statistically significant at 500 Hz (p < 0.1 %, Mann-Whitney-Rank-Sum-Test) and 1 kHz (p < 5%, Student t-Test). Beside the increase in obtained output amplitude the variability at low frequencies was decreased by the coupler / TUTOPATCH® combination. By using TUTOPATCH® and the coupler a major reduction the output variability (SD = 2.4 dB) compared to the “naked” FMT situation (SD = 9.2 dB) was found at 500 Hz

Conclusion
Applying the FMT with pre-defined force in combination with TUTOPATCH® and the RWC to the RW, leads to higher output and lower variability in sound transmission at low frequencies.
Fiber Arrangement in the Rat Tympanic Membrane
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Background
It is well known that the arrangement of fibers (radial and circular) in the pars tensa plays an important role in the mechanical properties of the tympanic membrane (TM). Scanning electron microscopy (SEM) images of the fiber arrangement of the TM have previously been reported. However, due to the limitations of available scanning electron microscopes at the time, those images were not able to resolve the detailed structure of the TM fiber network. Here we report the fiber arrangement of the rat TM observed under a high-resolution field emission SEM.

Methods
The TMs were dissected from adult rats and were fixed in 2.5% glutaraldehyde in phosphate buffer solution overnight. After decalcification in 5% EDTA for 24 hours, the TMs were soaked in 10% NaOH for 3-4 days and conductive-stained with 1% tannic acid followed by 1% OsO4. The specimens were then dehydrated in a series of graded ethanol solutions and critical point-dried using liquid CO2. The dried specimens were coated with 5 nm osmium and the fiber arrangement of the TM was observed under SEM from both the external ear canal side and the middle ear cavity side.

Results
The TM is composed of individual fibers with diameters around 20 nm. From the external ear canal side, radial and parabolic fibers can be observed. In the middle region between the manubrium and tympanic ring (TR), the orientation of the radial and parabolic fibers is highly ordered. Close to the TR, the radial fibers become intertwined into a random fiber network, and then attach to the TR. Similarly, the radial fibers become random near the manubrium and cover it. Observing from the middle ear cavity side, the radial fibers converge onto the manubrium in an orderly way. Circular fibers can be seen starting at a distance of one-tenth of the radius from the manubrium. They are woven into a plexus in the middle region and grouped into bundles close to the TR.

Conclusion
With state-of-the-art SEM available today, the fiber morphology and arrangement of the TM can be clearly observed.
Experimental Study of Effects of Opening Middle-Ear Cavity on Vibrations of Gerbil Tympanic Membrane

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Background
Opening the middle-ear air cavity strongly affects the behaviour of the middle ear. This study extends previous observations by examining the effects of different sizes of the opening, and the effects of the opening on vibrations at multiple points on the tympanic membrane and manubrium.

Methods
A laser Doppler vibrometer at the Montréal Children’s Hospital is used to measure the vibration at multiple locations on the tympanic membrane and manubrium. After making measurements with a closed cavity, the cavity is opened to varying extents, starting with a hole approximately 1 mm in diameter and ending with the largest possible opening. At each intermediate stage measurements are taken several times at the umbo, and at the final stage measurements are taken at multiple locations.

Results
Opening the cavity with even a small hole causes an increase in the low-frequency magnitude and a shift of the main middle-ear resonance to lower frequencies, and introduces an anti-resonance. When the opening is progressively widened the anti-resonance progressively moves from about 2 kHz to about 9.5 kHz. With a closed cavity, a normal flat pars flaccida causes a magnitude minimum in the manubrium and pars-tensa responses at a frequency between 400 and 700 Hz. This minimum becomes progressively more shallow as the opening is progressively enlarged. When the cavity is opened, the ratio of the displacement magnitude of the umbo to that of the short process changes little from its closed-cavity value. Although the low-frequency magnitudes of points on the pars tensa increase with even the smallest opening, they are practically unchanged when the opening is subsequently enlarged. Opening of the middle-ear cavity does not change the break-up frequency of the pars tensa.

Conclusion
Our multi-point measurements show that, although the middle-ear resonance moves to a lower frequency, the mode of vibration of the manubrium does not change appreciably when the cavity is opened. Opening the cavity introduces an anti-resonance which shifts to higher frequencies as the opening is enlarged. There is also a progressive diminishing of the effect of the pars flaccida on the responses. The presence or absence of the cavity has no effect on the break-up frequency of the pars tensa. These results have implications for both experimental and modelling studies.

Finite-Element Modelling of the Newborn Ear Canal and Middle Ear

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Background
There exist several techniques for early hearing screening and diagnosis but the available screening procedures cannot distinguish clearly between conductive and sensorineural hearing loss in newborns, and the results of available diagnostic tests in very young infants are difficult to interpret. Admittance measurements can help to detect conductive losses but do
not provide reliable results for newborns. The newborn ear is anatomically very different from the adult one; for example, the newborn’s canal wall is not yet fully ossified and has a much lower stiffness than that of the adult.

Methods
Finite-element models of the newborn ear canal and middle ear were developed and their responses were studied for frequencies up to 2000 Hz. Material properties were taken from previous measurements and estimates, and the sensitivities of the models to these different parameters were examined. The simulation results were validated through comparison with previous experimental measures.

Results
Simulations indicate that at frequencies up to 250 Hz the admittance of the canal wall is comparable to that of the middle ear in the newborn. Above 250 Hz the admittance of the canal wall remains almost constant. For the middle ear, however, there is a clearly defined resonance peak in the vicinity of 1100 Hz that produces an admittance much larger than that of the canal wall. Above this frequency, the middle-ear admittance gradually decreases, reaching a value approximately twice that of the canal wall at 2000 Hz. Overall, the middle-ear admittance only dominates that of the canal wall in a narrow band around the middle-ear resonance.

Conclusion
Our model predicts that at the conventional tympanometric frequency of 226 Hz the newborn canal-wall admittance is comparable to that of the middle ear, so the canal-wall admittance is not negligible as it is in adults. In addition, this model suggests that tympanometry with probe tones in the vicinity of the middle-ear resonance frequency, where the middle-ear admittance dominates, improves the chance that the measured admittance will give useful information about the middle ear.

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Negative Middle Ear Pressure versus Ear Canal Pressure Variations: Effects on Wideband Energy Absorbance Measurements in Humans
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Background
A recent study reported the effects of negative middle ear pressure (-MEP) and positive/negative ear canal pressure (±ECP) on the middle ear function using otoacoustic emissions (Sun, 2012). Wideband energy absorbance (EA) has been proposed to characterize the forward middle ear transfer function. It is determined by the ratio of absorbed to incident sound energy. This technique offers larger SNRs than otoacoustic emissions thus preventing underestimation by way of noise floor effects.

Methods
Data were collected from 34 adults with normal hearing using an Interacoustics wideband tympanometer. EA measurements were analyzed from 0.236 to 8 kHz with ECP swept from +200 to -300 daPa under normal and -MEP conditions. MEP was estimated by the bandpass tympanogram peak pressure (BTPP). Subjects produced -MEP by performing a Toynbee maneuver. Data were grouped into two MEP ranges: > -100 daPa (20 ears) and ≤ -100 daPa (23 ears). EA under normal MEP (at BTPP) was subtracted from EA with pressure manipulations to define four effects: (1) -MEP effect (EA under -MEP at the ambient pressure); (2) compensation effect (EA under -MEP at BTPP); and (3, 4) +ECP and -ECP effects (EA under normal MEP at ECP = ±(MEP), respectively).

Results
Under normal MEP (mean = -15 daPa), mean ambient EA was ~0.07 and ~0.09 lower than EA at BTPP for the two groups. Under -MEP for Group 1, EA decreased for frequencies below 2 kHz (~0.38 at 1 kHz) and increased (up to ~0.13) above 2 kHz. For Group 2, the transition frequency for EA decrease and increase moved upward to 4 kHz with a maximal reduction of ~0.43 and enhancement of ~0.19. Difference in EA between compensated and normal MEP conditions was small (~0.05). The EA change by +ECP and -MEP was similar. In contrast, -ECP resulted in less reduction (~0.05) for low frequencies and less enhancement (~0.1) for high frequencies.

Conclusion
Negative MEP substantially alters EA in a frequency-specific pattern. This effect is clinically important even in ears undergoing everyday MEP variations. These findings should be considered when establishing clinical norms. Application of an equivalent ECP in ears with -MEP effectively corrects the altered EA. This procedure may improve diagnosis of
middle ear diseases with coexistent MEP. The effects of +ECP and -MEP are comparable while the -ECP effect is smaller. Positive ECP may be utilized to research -MEP effects on middle ear physiology.
Assessment of tympanic membrane-coupled ossicular integrity on mechanical and physiologic responses in a middle ear effusion model

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Background
Middle ear effusions (MEE) arising as a result of otitis media or Eustachian tube dysfunction are characterized by impaired mobility of the tympanic membrane (TM) and ossicular chain often resulting in a conductive hearing loss. The contribution of diminished ossicular mobility to the hearing loss seen in MEE, however, remains to be further characterized. We hypothesized that if ossicular coupling is interrupted during a MEE, there would be a reduced umbo velocity magnitude ($[V_U]$) and an increased cochlear microphonic (CM) threshold.

Methods
$[V_U]$ and CM responses were measured in response to tone pips ranging in frequency from 0.25 – 8 kHz and in intensity from 10 - 100 dB in chinchilla ears (N=5). $[V_U]$ and CM thresholds were recorded in normal ears (baseline) and fluid-filled bullae (fluid) simulating a MEE (2.0cc, 0.5 Poise silicone oil). Responses were then observed in two additional conditions: 1) after disarticulating the incudostapedial joint in baseline ears (baseline without chain) and 2) after filling the disarticulated middle ear with fluid (fluid without chain).

Results
Results showed reduced $[V_U]$ at all frequencies in simulated MEE with the ossicular chain intact. With introduction of ossicular chain disruption, $[V_U]$ in the fluid without chain versus fluid condition decreased significantly at low and medium but not high frequencies (~4, 5, and 2 dB, respectively). Elevated CM thresholds were greatest in conditions without an intact chain (re: normal). A 6 dB increase in CM threshold was observed during the fluid without chain versus the baseline without chain state at low frequencies. The slope ($a$) of the linear regression relating $[V_U]$ and CM threshold changes shows proportional change with MEE, however, the proportionality is lost with disarticulation in baseline and fluid states ($a = 0.94, 0.07, and 0.69$ dB/dB, respectively).

Conclusion
During MEE with interrupted ossicular coupling 1) changes in $[V_U]$ are consistent with reduced summative TM input to umbo (re: fluid) and 2) changes in CM threshold suggest attenuated difference in sound pressure between the oval and round windows (re: baseline without chain). Moreover, CM and $[V_U]$ relations may be of potential clinical utility providing a sensitive test to differentiate MEE from MEE with ossicular pathology [supported by: NIH NIDCD R01-DC01155].

Development of cartilage conduction hearing aid (3) - Monosyllable intelligibility in the noisy condition-

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Background
In 2004, professor Hosoi found that clear sound can be heard when a transducer touches softly on an aural cartilage. From the basis for the idea, a trial model of the cartilage conduction hearing aid has been completed (Figure 1b). The ring-shaped fitting part attached to the transducer is put in the entrance of the external auditory canal, and the vibrated
cartilage around the fitting part transmit sound information. Since the cartilage conduction hearing aid does not occlude the external auditory canal, it can be used as an alternative to an open-fitting hearing aid. Our previous study reported that the vibration of aural cartilage generates sound in an external auditory canal especially in the low frequency range (< 3 kHz), and the amplified sound is hard to leak from the canal, regardless of the opening. So conversely, is the cartilage conduction hearing aid sensitive to inflow of environmental noise through the opening? To answer this question, monosyllable intelligibilities of open-fitting and cartilage conduction are compared in noisy conditions.

**Methods**
First, a sound pressure level (SPL) to present monosyllable is determined. Three subjects with normal hearing took speech audiogram tests in a silent condition. The presentation methods were open-fitting and cartilage conduction (Figure 1). In each presentation, the SPL to obtain maximum discrimination score (SPLmax) was measured for each subject, and the presentation SPL was determined by SPLmax + 10 dB.

Second, the speech audiogram tests were carried out presenting a white noise. A loudspeaker to present the white noise was put on the left side of the subject (the distance: 3 m). The SPL of noise changed in 55, 58, 61, 64, and 67 dB. The presentation methods of the monosyllables were open-fitting and cartilage conduction at the left ear, and the presentation SPL was constant. The unused right ear was occluded by an earplug.

**Results**
Figure 2 compared the monosyllable intelligibilities for the open-fitting and cartilage conduction presentations in the noisy conditions. The monosyllable intelligibility for the cartilage conduction was higher than that for the open-fitting.

**Conclusion**
Although the both open-fitting earplug and cartilage conduction transducer keeps the external auditory canal opened, the cartilage conduction transducer is more robust against the coming noise. The open-fitting earplug is hard to amplify sound in the low frequency range, while the cartilage conduction transducer does not leak the low frequency sound, which is important to discriminate speech.
Figure 1. Presentation methods

Figure 2. Monosyllable intelligibilities in the noisy conditions for each presentation method (error bar: standard deviation)
Investigating the Emergence of Binaural Processing Strategies after Surgical Correction of Congenital Unilateral Conductive Hearing Loss

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Background
Long-term improvement in performance on binaural processing tasks has not been observed after surgical correction of congenital unilateral conductive hearing loss (CUCHL). Previous investigations have evaluated subjects only over the short-term (< 6 months after surgery). However, animal studies investigating auditory development and research on human subjects receiving bilateral cochlear implants indicate that auditory plasticity may exist after critical periods have passed.

Methods
Subjects underwent surgery at least 1.5 years before free-field testing (range 1.5 – 7 years). Pure-tone audiometry was performed before and after surgery. Sound localization was tested while each listener sat in the center of an arc of 19 loudspeakers spaced 10° apart, positioned at 0° azimuth. A train of four bursts of pink noise at 50 dB sound pressure level (SPL) with a +/- 4 dB level rove was emitted from each loudspeaker, with twenty repetitions per location. Listeners indicated the perceived sound location on a touch-screen, and the difference between reported and actual locations was calculated as root mean square (RMS) error. Speech-in-noise testing involved listening to a male target voice consisting of CNC words presented from 0° azimuth. Two female maskers were presented at 50 dB SPL in the following configurations: 0°, +90°, -90°, or +/- 90°. Testing was conducted at signal-to-noise ratios (SNR) ranging from 0 to -30 dB SPL. Listeners were asked to select the heard word from a closed list of words. Spatial release from masking (SRM) was computed as the difference in performance with maskers at 0° vs. either +90°, -90°, or +/-90°.

Results
Preliminary results suggest that listeners demonstrate SRM within the range of values of normal-hearing controls (range 4-9 dB). Localization accuracy may be slightly worse in subjects tested to date (RMS range 8.37-12.99°) than in normal-hearing controls (range 5.29-9.2°), but is within the range of performance observed in normal-hearing children, and better than performance seen in bilateral cochlear implant users.

Conclusion
In listeners with CUCHL performance on binaural hearing tasks falls within the range observed in normal-hearing listeners, which has not been reported in patients with short-term follow-up. This may be a result of auditory plasticity. These findings suggest that binaural processing strategies may mature with continued auditory stimulation over long-term follow-up after repair of CUCHL.

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HB-EGF stimulates hyperplasia of the middle ear mucosa in bacterial otitis media
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Background
Otitis media is one of the most common pediatric diseases. While it is usually treated without difficulty, up to 20% of children may progress to long-term complications that include hearing loss, impaired speech and language development, academic underachievement, and irreversible disease. Hyperplasia of middle ear mucosa contributes to the sequelae of acute otitis media and is of important clinical significance. Understanding the role of growth factors in the mediation of mucosal hyperplasia could lead to the development of new therapeutic interventions for this disease and its sequelae.

Methods
To explore growth factors that may regulate mucosal hyperplasia, a whole genome gene array analysis of mRNA expression of mice during acute otitis media induced by nontypable Haemophilus influenzae was performed. We identified growth factors with expression kinetics temporally related to mucosal hyperplasia. We then tested the factors for their ability to stimulate mucosal growth in vitro with mucosal explant model of rat middle ear mucosa.

Results
From the gene array analysis, we discovered seven candidate growth factors with up-regulation of mRNA expression related to mucosal hyperplasia; Amphiregulin, Epiregulin, HB-EGF, GDF-15, LIF, Activin A, Cxcl1. Each factor was then applied in vitro to middle ear mucosal explants in three different concentrations. Of the seven, only HB-EGF induced significant mucosal hyperplasia. Subsequent quantification of HB-EGF protein expression in vivo via Western blot analysis with anti-HB-EGF antibody confirmed that the protein is highly expressed from 6 to 24 hours after bacterial inoculation, while immunohistochemistry revealed production by middle ear epithelial cells and infiltrated lymphocytes.

Conclusion
HB-EGF is expressed during the course of acute otitis media and can stimulate mucosal growth, suggesting an active role in the disease process. These results imply that therapies which target HB-EGF may have the ability to ameliorate mucosal hyperplasia during otitis media, and thereby reduce the detrimental sequelae of this childhood disease.

High-Throughput Screening of Kinases Regulating Mucosal Growth in Bacterial Otitis Media
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Background
Hyperplasia is a phenomenon unique to the mucosa of the middle ear. In the setting of otitis media it can contribute to the numerous detrimental sequelae of this common pediatric disease, including hearing loss, speech and language delay, and learning impediments. Understanding the mechanism of mucosal growth during infection is, therefore, of clinical significance. Though it is known that mucosal cell proliferation is regulated by intracellular signal transduction, the role of various signalling pathways in the middle ear remains poorly understood. We developed a novel high-throughput screening model to identify signaling cascades integral to mucosal hyperplasia, and used the model to evaluate the inhibition of kinase inhibitors.

Methods
In 48 well plates, we plated 0.25mm² explants of middle ear mucosa from bacterially infected rat ears. The mucosa was allowed 48 hours to attach. Kinase inhibitors from a 160 compound commercial library that covered the mammalian kinome were added to the culture media. Each inhibitor was applied at three concentrations to groups of three explants per concentration to develop a dose response function. The screening yielded compounds with the potential to regulate mucosal hyperplasia in bacterial otitis media. Elements of several different kinase families were found to contribute to mucosal hyperplasia.

Conclusion
We have established a high-throughput screening technique to identify kinase inhibitors that play a role in mucosal growth in the middle ear. This method proved to be repeatable and consistent, and may be useful in screening other compound
libraries in the future. Most importantly, it facilitates greater understanding of the intracellular signaling pathways involved in middle ear infection and introduces potential targets for the amelioration of otitis media sequelae.

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Development of cartilage conduction hearing aid (2) – benefits in the patients with continuous otorrhea or acquired aural atresia -
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Background
Sound was effectively transmitted by attaching a transducer to the aural cartilage even without fixation pressure. This new method for sound transmission was termed cartilage conduction (CC). CC can be utilized even in hearing-impaired patients who cannot use air-conduction hearing aids due to continuous otorrhea or aural atresia. Although we develop a hearing aid employing CC, the benefits of the prototype hearing aid have not yet been assessed. In this study, the CC hearing aid was fitted in patients, and its benefits were investigated.

Methods
Four patients with conditions such as continuous otorrhea and acquired aural atresia after surgery participated in this study. The benefits were assessed by audiometric tests and interview.

Results
Thresholds and speech recognition scores improved in all subjects. Below 2 kHz, the functional gains in the patients with acquired aural atresia were larger than those with continuous otorrhea. However, at 4 kHz, the functional gain at 4 kHz could not reach a sufficient level in the patients with aural atresia. The large gains below 2 kHz in the patients with aural atresia were probably caused by a soft tissue transmission pathway in the postoperative space. Sound at 4 kHz may be transmitted via direct air-conduction pathway. No subjects complained of pain associated with the attachment of the transducer, although such problems are usually observed for a bone-conduction hearing aid. This feature is considered one of the advantages of the CC hearing aid.

Conclusion
These results indicate that the CC hearing aid has potential as a useful amplification device for patients with hearing disability.
The effect of visual information on intelligibility in bone-conducted ultrasound perception

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Background
Since Gavreau reported that ultrasound could be heard by bone-conduction, many interesting characteristics have been reported about bone-conduced ultrasonic perception. Especially, recent studies indicated that Bone-conducted ultrasound (BCU) modulated by different speech sounds can be discriminated by some profoundly deaf subjects as well as the normal-hearing. Recently, a bone-conducted ultrasonic hearing-aid (BCUHA) for profoundly deaf individuals have been developed and its utility has been evaluated. However, previous studies using BCUHA revealed intelligibility only with the use of acoustic media in transmitting language information. In this study, we investigated the effects of visual information (lip-reading information) on intelligibility in bone-conducted ultrasound perception of normal-hearing individuals.

Methods
Speech discrimination tests were performed in three condition:(1) audio-alone condition, (2) audio-visual condition,(3) visual alone condition. The results of three speech discrimination tests were compared.

Results
The results of speech discrimination tests in this study revealed that lip-reading information strongly affected bone-conducted ultrasound speech perception. The speech intelligibility of bone-conducted ultrasound with visual information was significantly higher than that without visual information.

Conclusion
We found that lip-reading information had clear effects on bone-conducted ultrasound perception, showing that simultaneous presentation of audio and visual information improved intelligibility to levels sufficient for speech perception. Our findings also suggested the efficacy of use of signal processing techniques in improving the intelligibility of prior consonants.
Identification of Conductive Hearing Loss in CBA/CaJ Mice Using Bone Conduction
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Background
We explored the use of bone conduction to identify conductive hearing loss in mice.

Methods
Auditory brainstem responses (ABR) were recorded from two dozen 7 to 8-week old CBA mice with both acoustic and bone conduction stimuli. For bone conduction stimulation, a flat head metal screw was implanted near the vertex of the skull, and the screw was glued to a Bruel & Kjaer minishaker. Brief tone pip stimuli were presented either by air conduction (AC) or by activating the shaker (BC). ABR thresholds, the lowest stimulus levels to produce response peaks, were determined by visual detection. Conductive hearing pathology was introduced by two ways: 1. removal of the tympanic membrane (TM) and malleus of the tested ear, 2. filling the middle ear cavity (MEC) with saline. Removal of the TM and malleus decouples the ossicular chain that is required to efficiently transfer acoustic energy into the cochlea. With the fluid filled middle ear cavity, the middle system now becomes stiffer making it harder for the TM and ossicles to move in response to AC sound.

Results
Both of these manipulations produced significant increases (30-50 dB) in ABR thresholds in response to acoustic stimulation. Since bone conduction involves different pathways, we expected that these manipulations would have little effect on the hearing sensitivity. Our results show that the threshold stimulus levels with bone conduction stimulation remained unchanged at frequencies above 10 kHz, and at lower frequencies, were increased by 5-10 dB.

Conclusion
These results suggest that the combination of air and bone conduction testing can distinguish sensorineural hearing loss from conductive hearing loss in mice. This technique can be a useful tool in evaluating the onset and development of hearing loss in various strains of mice.

Finding of Characteristic DFNA9 Cochlin-staining Eosinophilic Deposits in the Middle Ear: Comparison of Wild-type (WT), Coch Knock-In (KI) and Knock-Out (KO) Mouse Models and DFNA9-Affected and Unaffected Individuals
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Background
Mutations in COCH, encoding cochlin, the most abundant protein detected in the inner ear, cause the midlife-onset, progressive sensorineural hearing loss and vestibular disorder, DFNA9. To date, 14 missense COCH mutations have been found throughout four continents. A pathognomonic DFNA9 finding is presence of cochlin-staining, eosinophilic deposits throughout the spiral ligament and limbus, and the stroma in the vestibular system, and reduction in number of fibrocytes, which normally express cochlin. A recent study (McCall AA et al., JARO, 2011, 12(2):141-9) reported presence of acellular deposits in middle ear structures in DFNA9. We have investigated more thoroughly the nature of these deposits and the role of cochlin by characterizing 1) middle ear histology in wild-type (WT), CochG88E/G88E knock-in (KI) and CochG88E/G88E knock-out (KO) mouse models, 2) cochlin localization in unaffected human and WT mouse middle ears, and 3) cochlin immunostaining in human DFNA9 and Coch KI mouse middle ears.

Methods
H&E staining of celloidin-embedded DFNA9 and age-matched control middle ear sections, and paraffin-embedded WT, Coch KI, and Coch KO mouse sections were performed. Immunohistochemistry of adjacent sections were performed using an anti-cochlin antibody.

Results
Eosinophilic deposits were detected in the Coch KI middle ears, most notably in the inter-ossicular joints. These aggregates were absent in the WT and Coch KO mouse, consistent with our hypothesis that DFNA9 pathology is gain of deleterious function of mutant cochlin, rather than haploinsufficiency, or absence of cochlin. Immunostaining revealed specific and prominent cochlin localization in inter-ossicular joints and pars flaccida of the tympanic membrane (TM) in both WT and Coch KI mice, including areas of aggregates. Deposit formation in the TM of 1-year-old KI mice was not
appreciable, in contrast to prominent deposits in the incudo-malleal joint, suggesting that abnormal aggregates in TM occur later than in the inter-ossicular joints. Mice at different ages are being evaluated to assess initial formation and progression of this pathology. Immunostaining of DFNA9-affected and age-matched control middle ears revealed cochlin localization in the same structures as seen in the mouse, as well as in the aggregates. Proteomic studies underway may further identify composition of the aggregates, pointing toward cochlin interactors and other proteins involved in cochlin functional pathways.

Conclusion
Our DFNA9 and Coch KI findings provide further insight into cochlin function and its role in progression of the disease process.

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Therapeutic Potential of Adenovirus-mediated Induction of β-defensin 2 for Otitis Media
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Background
In our prior studies, we demonstrated that β-defensins inhibit viability of the common otitis media pathogens. We also showed that the middle ear epithelial cells induce β-defensin 2 in response to nontypeable H. influenzae (NTHi) via TLR2-dependent signaling, suggesting therapeutic potential of a β-defensin 2 inducer. Here, we aim to determine if an adenoviral vector containing the human β-defensin 2 gene (DEFB4) can serve as a β-defensin 2 inducer of the middle ear epithelial cells.

Methods
Regulation of β-defensin 2 expression was determined with qRT-PCR analysis and ELISA analysis. Confocal microscopy was performed to determine an inhibitory effect of β-defensin 2 on bacterial adhesion. For bacterial clearance, NTHi colonies were counted after culturing of the murine middle ear lavage.

Results
We found that transtympanic inoculation of the recombinant β-defensin 2 enhances NTHi clearance from the middle ear cavity. The adenoviral vector expressing β-defensin 2 (Ad-DEFB4) was found to up-regulate β-defensin 2 expression in the epithelial cells in a time- and dose-dependent manner, compared to Ad-null. Infection with Ad-DEFB4 appeared to inhibit adhesion of NTHi to the epithelial cells. Transtympanic inoculation of Ad-β-gal appeared to induce β-galactosidase without inflammatory response in the rat middle ear. Moreover, Ad-DEFB4 was found to improve clearance of NTHi from the murine middle ear cavity.

Conclusion
Taken together, our results suggest that induction of β-defensin 2 by an adenoviral vector may be considered as an alternative strategy for the management of middle ear infection. [Supported by NIH grants: DC005025, DC006276 and DC011862]
Influence of target tissue on survival and differentiation of neurons derived from hES cells
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Background
Cell replacement therapy is a potentially powerful approach to replace degenerated or severely damaged hair cells and spiral ganglion neurons in the inner ear. Cell therapy has been proposed to be a promising strategy to restore hearing either by replacing degenerated neurons or via improving the efficacy of cochlear implants which rely on functional neurons. Neurons derived from embryonic and adult neural tissue or stem cells have been successfully transplanted into the inner ear. It remains to identify how the auditory tissue influence these implanted cells. During development and regeneration, a plethora of growth factors are secreted from the target tissue including neurotrophins. These factors promote the survival, migration and integration of the approaching cells. Currently, we are investigating how the secreted factors from the auditory tissue influence the growth and differentiation of neural progenitors generated from human embryonic stem (hES) cells.

Methods
Organotypic culture from different tissue (brainstem, organ of Corti, spiral ganglion neurons) were prepared from early postnatal rat. Conditioned media which contains all the secreted factors were collected. The activity of the medium from the different auditory system tissues was tested on neurons derived from human neuroblastoma SHSY5Y or hES cells cultures. Different parameters using molecular biology, immunohistochemistry and cytotoxicity and other techniques are tested.

Results
The preliminary results of the testing of the effects of secreted factors on the neural progenitor derived from hES cells exhibit interesting trends. The measurement of the cell migration distance in these progenitors shows that medium from SGNs and organ of Corti induced higher migration while brainstem derived medium was similar to control. The neurite outgrowth is parameter to check the differentiation of progenitor into neurons. Again we observed that the medium from SGN induce higher neurite outgrowth in the hES cells derived neural progenitors then control.

Conclusion
Our preliminary data shows that there are factors secreted from the auditory tissue which influence the differentiation of the neurons derived from hES cells. Identification of these factors would be very helpful in developing cell therapy for inner ear.
Localization of kainate receptors in inner and outer hair cell synapses
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Background
Glutamate and its receptors mediate inner hair cell (IHC) afferent transmission. Recent physiological data show that glutamate also plays a role in outer hair cell (OHC)/type II afferent transmission, but the glutamate receptors that are involved are not well defined. A previous study using in situ hybridization showed that several kainate-type glutamate receptor subunits (GluR5, GluR6, KA1, KA2) are expressed in cochlea ganglion neurons (Niedzielski & Wenthold, 1995, J Neurosci 15: 2338). Another group reported that kainate receptors are expressed in IHC afferent synapses and contribute to hair cell acoustic transmission (Peppi et al., 2012, J Assoc Res Otolaryngol 13: 199).

Methods
X-gal staining was performed on whole-mount cochlea and cryostat sections of KA2-KO mice (P8, P14, 1.5 months and 1 year old), which have a Lac Z cassette for the reporter gene. Type I and type II ganglion neurons were discriminated by different labeling patterns of neurofilament H. Immunocytochemistry was performed on rat whole-mount cochlea using commercial antibodies against all kainate receptor subunits. Slight paraformaldehyde fixation combined with microwave-mediated antigen-retrieval or pure methanol fixation was essential for labeling. Also, electron microscopy of postembedding immunogold labeling was performed in guinea pigs.

Results
X-gal staining was found in both type I and type II ganglion neurons in adults. Also, staining was seen only in OHCs and spiral limbus region at 1.5 months or older, although staining may be more widespread at younger ages.

GluR6 immunolabeling colocalized with CtBP2 at IHC and OHC synapses. Also, labeling was contiguous with postsynaptic PSD-93 at OHC synapses; these patterns matched those with the AMPA glutamate receptor subunit, GluR2. GluR5 mainly labeled OHC efferent synapses - shown by double labeling of VAMP2 or Na⁺/K⁺-ATPase α3. No distinct labeling was found in IHC and OHC afferent synapses. KA2 labeled IHC afferent synapses definitively, but labeling in OHC synapses was less definitive. KA1 appeared to label OHC afferent synapses but not IHC ones. No immunoreactivity for GluR7 was detected in the cochlea.

Conclusion
OHC afferent synapses contain at least GluR6 and possibly KA1 and KA2 that could form various combinations on the presynaptic and postsynaptic membrane. GluR6 and KA2 are expressed in IHC afferent synapses. These results suggest that kainate receptors are involved in synaptic transmission in both IHCs and OHCs. Also, GluR5-containing receptors may modulate OHC efferent inhibition.

Exocytosis of protons blocks Ca²⁺ currents in auditory hair cells
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Background
Ca²⁺ influx across the plasma membrane of hair cells triggers neurotransmitter release at ribbon type active zones. To manage an uninterrupted flow of transmission, this synapse continuously releases glutamate, the neurotransmitter stored in synaptic vesicles. The pH inside vesicles is thought to be about 5.7 and the exchange of protons for glutamate allows the vesicular glutamate transporter to fill synaptic vesicles with glutamate. The exocytosis of synaptic vesicles transiently acidifies the synaptic cleft of bipolar cell synapses, which are also ribbon type synapses. This leads to a transient acidification of the synaptic cleft and block of the Ca²⁺ channels located near the sites of exocytosis. We studied an analogous process in the hair cell ribbon synapse.

Methods
Amphibian papillae were carefully dissected from adult bullfrogs that had been sedated in ice water bath and double-pithed. Semi-intact preparations of hair cells and their connecting afferent fibers were obtained as described by previous studies. Whole-cell patch-clamp recordings were performed with an EPC-10/2 patch-clamp amplifier (HEKA).

Results
We have found that the shapes of Ca²⁺ currents are different depending on the membrane holding potential of hair cells. When hair cells were depolarized from -60 mV, a potential close to the physiological resting membrane potential of in vivo hair cells, to -30 mV, the Ca²⁺ currents showed “outward” transient components in contrast with Ca²⁺ currents of hair cells
that were held at -90 mV. The average size of outward component was 22.9 ± 2.6% of the peak Ca\(^{2+}\) current. We also examined how much time is required to for a detectable outward component to develop. We depolarized hair cells from -90 mV to -60 mV for various durations as a pre-pulse and then depolarized to -30 mV for 20 ms. The outward component required about 50 ms to 100 ms at -60 mV to start showing, and the size of this component increased as hair cells were held longer at -60 mV. To confirm whether protons co-released with glutamate from fused vesicles cause the outward components in Ca\(^{2+}\) currents, we applied methylamine, a weak base. This abolished the outward component. The outward component was also significantly decreased when the concentration of external HEPES was increased to 50 mM.

**Conclusion**

The transient outward components in Ca\(^{2+}\) currents of frog auditory hair cells results from a block of Ca\(^{2+}\) channels by protons released from fused synaptic vesicles.

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**Evidence for K+ accumulation around mammalian vestibular Type I hair cells: an electrophysiological study**

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**Background**

Mammalian vestibular organs contain two types of sensory receptors, named Type I and Type II hair cells. While Type II hair cells are contacted by several small afferent nerve terminals, the basolateral surface of Type I hair cells is almost entirely enveloped by a single large afferent nerve terminal, called calyx. Moreover Type I, but not Type II hair cells, express a low-voltage activated outward K\(^+\) current, IK,L, which is responsible for their much lower input resistance (Rm) at rest as compared to Type II hair cells. The functional meaning of IK,L and associated calyx is still enigmatic.

**Methods**

Experiments were performed on mice (C57 and Swiss CD1) obtained from Harlan Italy. Mouse age ranged between postnatal day (P) 7 and P25. The roof of the isolated SCC ampulla was gently torn to expose the crista ampullaris, which was then immobilized at the bottom of a glass Petri dish by means of a weighted nylon mesh. Hair cells in the crista were viewed and photographed by using an upright microscope equipped with differential interference contrast optics (Zeiss Axioskop, Germany), 63x water immersion objective. By using the patch-clamp whole-cell technique we have recorded the current- and voltage responses from in situ hair cells.

**Results**

Outward K\(^+\) current activation resulted in K\(^+\) accumulation around Type I hair cells, since it induced a rightward shift of the K\(^+\) reversal potential, the magnitude of which depended on the amplitude and duration of K\(^+\) current flow. Since this phenomenon was never observed for Type II hair cells, we ascribed it to the presence of a residual calyx limiting K\(^+\) efflux from the synaptic cleft. Intercellular K\(^+\) accumulation added a slow (tau > 100 ms) depolarizing component to the cell voltage response. In a few cases we were able to record from the calyx and found evidence for intercellular K\(^+\) accumulation as well. The resulting depolarization could trigger a discharge of action potentials in the afferent nerve fiber.

**Conclusion**

Our results support a model where pre- and postsynaptic depolarization produced by intercellular K\(^+\) accumulation cooperates with the conventional neurotransmitter release in sustaining afferent transmission arising from Type I hair cells. While vesicular transmission together with the low Rm of Type I hair cells appears best suited for signaling fast head movements, depolarization produced by intercellular K\(^+\) accumulation could enhance signal transmission during slow head movements.

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Spatio-temporal pattern of action potential firing in mouse inner hair cells
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¹Inserm

Background
Inner hair cells (IHCs) are the primary transducer for sound encoding in the cochlea. In contrast to the graded receptor potential of adult IHCs, immature hair cells fire spontaneous calcium action potentials during the first postnatal week.

Methods
Here, we investigate the pattern of spontaneous action potential firing in mammalian IHCs during development. We recorded action potentials from neonatal mouse IHCs (P1 to P7) of apical or basal cochlear coils using the perforated patch-clamp technique in the current-clamp mode (|inj|=0pA).

Results
We show that IHCs fire burst of action potential and that pattern is undistinguishable between basal and apical hair cells. However, the bursting behavior became more salient during development, i.e, from a broad structure right after birth to a sharp and stereotyped motif at the end of the first postnatal week.

Conclusion
The IHCs spiking activity has been proposed to shape the tonotopic map along the ascending auditory pathway. We suggest that in addition to carrying place information to the ascending auditory nuclei, the firing pattern conveys temporal information of the cochlear development.

Analysis of Fluorescently-Labeled Protein Puncta in Hair Cells: Synaptic Strength & Systematic Artifacts
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Background
Immuno-fluorescence microscopy promises to provide high-throughput assessment of antigen abundance in the organ of Corti, at the level of individual synapses. However, semi-quantitative measurements of protein levels may be significantly biased by artifacts introduced during image acquisition. We estimated and corrected for artifacts of (1) bleaching and (2) attenuation in the optical axis so that molecular determinants of relative synaptic strength can be ascertained with confidence.

Methods
Synapses were triple-labeled for glutamate receptors, calcium channels, and synaptic ribbons. (1) The effect of bleaching was assessed by acquiring two Z-stacks on the same volume of tissue and changing the order-of-acquisition in Z, from top-down to bottom-up. (2) The effect of attenuation was assessed by mounting the organ between two cover slips and imaging the same region of interest from both sides. Puncta were identified by thresholding and then quantified by integrating the voxel intensities within iso-intensity surfaces after background subtraction. Standard microscope settings: 10% laser power, 0.33µm step size, 4 scans averaged per XY-section, 400Hz scan rate, 50nm pixels, 25x50µm field of view).

Results
(1) The rate of bleaching varied amongst the 3 labels in a stack, ranging from 1-37% when measured as the mean reduction of punctum intensity between two 20µm Z-stacks acquired one after another on the same volume (20x3x4=240 XY-scans per channel per stack). Thus the order in which a Z-stack was acquired resulted in as much as ~37% difference in measured intensity of an individual punctum, depending upon whether it was imaged in the first or the final XY-section of the stack. (2) Signal attenuation as a function of tissue depth produced a systematic intensity gradient in the optical axis when using conventional mounting media. When a special mounting medium (TDE) was used to match the refractive index of tissue, attenuation and the resulting spatial gradient artifact were relatively small.

Conclusion
Intensity correlations between labels and spatial gradients of label intensity can be artificially created, augmented, or obscured by systematic artifacts of image acquisition. They can be averaged-out by varying the order-of-acquisition and the preparation’s orientation. Before correlations can be interpreted and cells reconstructed with cell-centric coordinates,
these Z-axis artifacts must be corrected for. We present a strategy for estimation and compensation to minimize these artifacts of bleaching and attenuation.
Evolutionary changes of the efferent hair cell receptor

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Background
The α9 and α10 nicotinic acetylcholine receptor (nAChR) subunits are expressed in cochlear hair cells, where they form the receptor that participates in the efferent control of auditory function. This nAChR mediates the inhibitory synapse between efferent fibers and outer hair cells in mammals, or short hair cells in birds. This cholinergic efferent feedback to hair cells is a common feature among all vertebrates. Accordingly, one would expect the evolutionary history of the genes coding for the α9 and α10 subunits to be similar throughout the whole Subphylum. However, a detailed analysis of the phylogeny and sequences showed signatures of positive selective pressure only for the mammalian α10 subunits. These differences in the evolutionary history correlate with the divergent biophysical properties of rat and chicken α9α10 nAChRs, such as calcium permeability. Here, we compared the pharmacology of chicken and rat receptors looking for a better understanding of the effects of the non-synonymous substitutions accumulated on mammalian α10 subunits.

Methods
Recombinant α9 and α10 subunit cRNAs were synthesized in vitro and injected in Xenopus laevis oocytes. Responses to drugs were measured using the two microelectrode voltage-clamp technique.

Results
The effects of the classical nicotinic agonists choline and DMPP and the antagonist serotonin on chicken receptors differed from those on rat α9α10 receptors and resembled those on rat α9. Above all, choline showed higher efficacy on chicken α9α10 receptors (88±8% max response to ACh, n=5) compared to rat α9α10 receptors (34±4% max response to ACh, n=4). Moreover, in a Ratα9Chickα10 hybrid receptor, responses to choline were 86±4% (n=3) of the maximal response to ACh, resembling chicken receptors and strongly suggesting that the sites involved have been altered in mammalian α10 subunits.

Conclusion
The aminoacid changes that accumulated on mammalian α10 subunits resulted in a mammalian α9α10 nAChR with reduced efficacy to the neurotransmitter metabolite choline. We propose that differential sensitivity to choline might result in differences in efferent synaptic kinetics and strength across species.
Miniature Post Synaptic Currents are Entrained by Infrared Pulses
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Background
Inner ear hair cells are exquisitely sensitive to pulsed infrared radiation (IR) and often modulate neurotransmitter release in response to each optical pulse. Four distinct endogenous mechanisms of IR sensitivity have been demonstrated: optoacoustic gating of the mechano-electrical transduction (MET) channels, thermal modulation of plasma membrane ion channels, thermal modulation of plasma membrane capacitance, and thermal modulation of intracellular Ca$^{2+}$ signaling. In the present work we examined thermal modulation of synaptic vesicle release by single IR pulses at wavelengths where IR absorption by water converts the IR optical energy into rapid, but small, heat pulses.

Methods
The neuro-muscular junction of Caenorhabditis elegans was used as the model system to facilitate genetic manipulations and reveal molecular targets of IR energy. Conventional whole cell voltage clamp was used to record spontaneous and laser-pulse evoked miniature post-synaptic currents (mPSCs). IR pulses were delivered via a 200-400 µm diameter optical fiber positioned approximately one fiber diameter away from the cell. Laser diodes tuned to 1862nm or 1550nm were gated at 0-500 µs pulse to deliver 0-20 µJ/pulse over and area of 100-300nm$^2$.

Results
Post-laser-stimulus histograms revealed momentary silencing of mPSCs (zero probability at 0-1.5ms) followed by a doubling of the mPSC rate (peak probability ~2x spontaneous at ~2.6ms). Recovery to the background mPSC rate followed two time courses, a rapid component ~1ms accounting for nearly half of the peak response, followed by a slow component correlating with thermal relaxation. Increased mPSC rates occurred in zero extracellular Ca$^{2+}$ as well as calcium channel knock outs (nca-1; nca-2) showing that voltage activated plasma membrane Ca$^{2+}$ currents were not required and that IR acted primarily through a voltage independent synaptic vesicle release mechanism.

Conclusion
Pulsed IR rapidly alters the capacitance of the membrane through charge redistribution, and evokes mitochondrial Ca$^{2+}$ currents through non-equilibrium thermodynamics. Although causality has not been shown, present results are consistent with the hypothesis that these biophysical events underlie IR entrainment of mPSCs reported here and IR-evoked synaptic transmission reported previously in vestibular hair cells.

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Protocadherin-15 (PCDH15) interacts with integrin α8 (Itga8) in the inner ear. Implications for a signaling mechanism involving Usher proteins.
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Background
In the inner ear PCDH15 is expressed at the apical and basal aspects of hair cells and neuronal terminals, with multiples isoforms playing different roles, from the kinociliary and tip links formation, to cell polarity and synapse maturation. Mutations in the PCDH15 gene result in Usher syndrome type 1F (USH1F) and sometimes non-syndromic deafness. Itga8 is an integral membrane protein that, upon dimerization with integrin β1, forms an active membrane receptor that binds extracellular matrix ligands and signals mainly through the small GTPase, RhoA. Expression of Itga8 has been demonstrated in the mouse apical hair cell surface and a possible association with focal adhesion kinase has been suggested.

Methods
We employed dual confocal immunohistochemistry, siRNA knockdown technology in the mouse embryonic hair cell line UB/OC-1, and co-immunoprecipitation studies to address whether a possible association between itga8 and PCDH15 exists.

Results
We observed basal and apical expression of itga8 in P1 mouse cochlea. In zebrafish ITGA8 is expressed at the apical and basal aspects of neuromast hair cells, where it co-localizes with Pcdh15a and Pcdh15b and the pre-synaptic marker...
SV2. UB/OC-1 cells transfected with siRNAs directed to itga8 showed a modification in the distribution pattern for PCDH15 compared with the scrambled controls, suggesting a key role for itga8 in PCDH15 trafficking and/or protein stabilization. PCDH15 knockdown cells did not show any effect in itga8 expression or distribution. In agreement with this last result, the expression analysis in the PCDH15 mutant mouse model, av-3J, show the characteristic distribution pattern and protein abundance for ITGA8 compare with wild type animals. These results reinforce the idea of itga8 playing a central role in complex formation. Co-immunoprecipitation studies from P1 mouse inner ear demonstrated the existence of this complex in vivo.

Conclusion
Our results demonstrate protein interactions between specific PCDH15 variants and itga8. This is the first evidence linking Usher protein function to an integrin-mediated signaling cascade. Because both proteins are expressed early in the immature inner ear they may function in hair cell stereocilia and synaptic maturation. Their expression in zebrafish during hair cell development provides evidence of a highly conserved function for these two proteins.
This work was supported by NIH grants R01 DC004844 and 5 P20 RR018788-08.

Stereocilia Tenting is Present in Live Hair Cells and Involves Remodeling of the Actin Core
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Background
Tenting refers to the “conical” or “wedge-like” shape at the stereocilia tips of the auditory and vestibular hair cells. It has been repeatedly imaged using scanning (SEM) and transmission (TEM) electron microscopy. Some TEM images show actin filaments filling the tented stereocilia tips while other TEM images depict membrane tenting only. The prevailing model proposes that tenting is caused by the tension that tip links exert on the plasma membrane. This model is mainly supported by SEM data of chemically-fixed tissue showing the loss of tenting after the disruption of tip links via chemical or genetic manipulations. However, specimen preparation artifacts (such as tissue shrinkage) could result in an exaggerated tented appearance of the stereocilia when the tip links are present.

Methods
We imaged the stereocilia bundles of live rat inner hair cells using hopping probe scanning ion conductance microscopy (HPSICM). Using conventional SEM, we studied the dynamic changes in stereocilia tenting after the disruption of tip links with BAPTA-buffered Ca²⁺-free medium.

Results
HPSICM imaging showed, for the first time in live cells, the presence of tenting in the second and third rows of inner hair cell stereocilia. We were not able to find any differences between the shape of the stereocilia tips observed with HPSICM and SEM. Using SEM, we observed that the degree of tenting is different in outer versus inner hair cells. Moreover, the degree of tenting in inner hair cells showed a gradual increase from base to apex. Immediately after BAPTA treatment the tip links were not observed but, interestingly, tenting was still present. Tenting did disappear by 20 min post-BAPTA and the time course of its recovery correlated with the re-appearance of the tip links during the following 5 hours.

Conclusion
Our results demonstrate that stereocilia tenting is not an artifact of tissue fixation but is likely to be a tip link-mediated and mechanically-driven phenomenon that involves the remodeling of the stereocilia actin core.

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Eps8L2 knockout mice show progressive hearing loss and hair bundle defects in auditory hair cells

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Background
The mammalian auditory system is able to detect and process a wide range of sound frequencies and intensities. Central to this process is the transduction of sound waves into electrical signals by cochlear hair cells, which depends upon mechanosensitive stereociliary bundles (Fettiplace and Hackney, 2000 Nat Rev Neurosci 7: 19-29). The growth and maintenance of the stereociliary actin core is dynamically regulated via a treadmill mechanism with actin polymerization at the tip and depolymerisation at the base (Manor and Kachar 2008 Semin Cell Dev Biol 19: 502-510). Actin capping (e.g. twinfilin 2, gelsolin, Eps8) and cross-linking (e.g. espin, plastin) proteins are crucial for the elongation and thickening of the stereocilia actin core.

Methods
We used a combination of immunolabelling, overexpression of GFT-tagged proteins, single-cell electrophysiology, electron microscopy and in vivo physiology to investigate the role of Eps8L2, an Eps8-related protein endowed with actin binding activity (Offenhauser et al. 2004 Mol Biol Cell 15: 91-98).

Results
Using control and Eps8L2 knockout mice we show that this protein is a novel component of the hair-cell hair bundle. Eps8L2 localized at the tip of the stereocilia of hair cells from the cochlear and vestibular system and is essential for their maintenance. Eps8L2 knockout mice exhibited progressive hearing loss, which was directly linked to a gradual deterioration of hair-bundle morphology. We found that Eps8-L2 regulates the maintenance of stereocilia height and thickness in hair cells.

Conclusion
Our results indicate that Eps8-L2 directly regulates actin dynamics in hair-cell stereocilia and that it is critical in maintaining the normal functionality of hair cells and hearing in mammals.

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Mechanotransduction Defects in Sensory Hair Cells of USH1C Knock-In Mice.

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Background
Numerous mouse models of Usher Syndrome (USH) have been identified or engineered over the past decade. Interestingly from over seven models that target harmonin (USH1C), only one reproduces both auditory and retinal deficits. The Ush1c.216G>A (Lentz et al. 2007, 2009) is a knock-in model with the cryptic splice site mutation found in French-Acadian USH1C patients. The mutation results in the complete absence of ABR responses in homozygous mutant mice at one month indicating the mice are deaf. Cochlear histology at P30 shows disorganized hair cell rows, abnormal bundles, and loss of both inner and outer hair cells along the organ (Lentz et al. 2012).

Methods
To assess the development of hair cells in the Ush1c.216G>A mutant mouse, we analyzed hair bundle morphology, the developmental expression of voltage dependent channels as well as the presence of functional mechanosensitive ion channels in outer hair cells (OHCs) of wild type, heterozygous and homozygous littermates during the first postnatal week.

Results
Disorganized hair bundles were evident in homozygous mice along the entire organ of Corti. Normal voltage-dependent currents and resting potentials were observed for all genotypes. Analysis of FM1-43 uptake in control and heterozygous mutant mice revealed a gradual developmental increase base to apex during the first postnatal week as described
previously (Lelli et al. 2009). While the developmental pattern was maintained in the organ of homozygous mutant mice, weaker FM1-43 fluorescence was observed with some sensory cells showing strong dye uptake and others showing no uptake. Direct measurement of transduction currents evoked by stiff-probe deflections of OHC bundles mice revealed similar properties in heterozygous Ush1c.216G>A and wild type littermates with maximal currents (Imax) of 468 ± 53pA (n=8 het +/- S.D., Apical half P4-P6) at a holding potential of -64mV. In contrast but consistent with the decreased FM1-43 fluorescence, OHCs from homozygous mutants with moderately disrupted hair bundles had smaller (range: 66pA to 296pA) or no transduction current with a mean Imax of 220 ± 70pA (n=9 cells excluding non-transducing cells, Apical half, P3-P6). Adaptation was always present but with a slower fast component in homozygous mutants, similar to that described for other harmonin mutants. More complete adaptation was also occasionally observed.

Conclusion
We conclude that mechanosensitivity is partially preserved in developing OHCs of Ush1c.216G>A mutant mice.

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Localization of PDZD7 to the stereocilia ankle-link associates this scaffolding protein with the Usher syndrome protein network
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Background
Usher syndrome is the leading cause of genetic deaf-blindness. Monoallelic mutations in PDZD7 increase the severity of Usher type II syndrome caused by mutations in USH2A and GPR98, which respectively encode for plasma membrane proteins usherin and GPR98. PDZ domain-containing 7 protein (PDZD7) is a paralog of the scaffolding proteins harmonin and whirlin, which are implicated in Usher type 1 and 2 syndromes, respectively. While usherin and GPR98 have been reported to form ankle links in hair-cell stereocilia, harmonin localizes to the stereocilia upper tip-link density and whirlin localizes to both the stereocilia tip and ankle-link regions. Moreover, PDZD7 has been shown to co-immunoprecipitate with USH1 proteins that make up the upper tip-link insertion density, i.e., harmonin (USH1C) and sans (USH1G), as well as with the cytosolic tails of usherin and GPR98. These interactions have not been corroborated by structural evidence. We wanted to investigate the subcellular localization of PDZD7 in hair cells, and to understand its potential role in the development and function of their mechanosensory stereocilia.

Methods
We used mass spectrometry to examine the presence of PDZD7 amongst chicken purified hair bundle proteins. We used immunopurified PDZD7 antibodies to examine its subcellular localization in mammalian sensory hair cells stereocilia bundle and compare it to that of USH1 and USH2 proteins complexes. We also used ballistic transfection of explant rat hair cells to examine the subcellular localization of Cherry-tagged PDZD7. Finally, we used immunocytochemistry on LLC-PK1 epithelial cells to investigate the interactions of PDZD7 with proteins of the ankle-link complex.

Results
We show that PDZD7 is expressed in chick stereocilia at a comparable molecular abundance to GPR98. We also show by immunofluorescence and by overexpression of tagged proteins in rat hair cells that PDZD7 localizes to the ankle-link region of developing organ of Corti and vestibular hair cells and of only mature vestibular hair cells, overlapping with usherin, whirlin, and GPR98. Finally, we expressed in LLC-PK1 cells fusions of the cytosolic domains of both usherin and GPR98 and the interleukin-alpha subunit, which targets the plasma membrane, and show that both whirlin and PDZD7 can bind to both tails.

Conclusion
Our observations are consistent with PDZD7 being a modifier and candidate gene for USH2, and suggest that PDZD7 is a second scaffolding component of the ankle-link complex that positions and stabilizes GPR98 at hair cells ankle-link region.
Can side links mediate hair cell mechanotransduction?
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Background
The tip links are thought to be the tethers that open the mechanotransduction channels on the tops of the shorter row stereocilia. Two molecular components of the tip link have been identified: protocadherin 15 (PCDH15) at the lower end of the link and cadherin 23 (CDH23) at the upper end of the link. Both PCDH15 and CDH23 have been shown to be necessary for mechanotransduction in hair cells.

The asymmetric PCDH15/CDH23 composition of the tip link may be disrupted in Shaker 2 and Whirler mice. The inner hair cell bundles in these mutant mice do not form a characteristic staircase. Instead, their stereocilia are interconnected with “top-to-top” links that still mediate mechanotransduction responses but in both positive and negative directions.

Methods
PCDH15 and CDH23 were immune-localized using immuno-gold scanning electron microscopy (Immuno-SEM). The directional sensitivity of the mechanotransduction responses was studied with conventional whole cell patch clamp technique.

Results
In Shaker 2 and Whirler inner hair cells, stereocilia are packed in a hexagonal pattern with the links running in 6 predominant directions from each stereocilium, along 3 axes. We found that the links connecting stereocilia aligned in the axis of normal mechanosensitivity (presumably tip link equivalent) have the same amount of PCDH15 and CDH23 as the side links between neighboring stereocilia.

Conclusion
Our data suggests an apparently identical molecular composition of the tip link equivalent and side links in Shaker 2 and Whirler inner hair cells. Therefore it is very likely that side links could mediate similar mechanotransduction responses. The experiments studying the directionality of mechanotransduction responses in these bundles are underway.
Weak Lateral Coupling between Stereocilia of Inner and Outer Hair-Cell Bundles
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Background
The forces felt by different transduction channels in a bundle depend critically on how well stereocilia remain cohesive during deflection. In the bullfrog saccule, sliding adhesion mediated by horizontal top connectors confers coherent motion to hair cell stereocilia and parallel gating to all transduction channels (Karavitaki and Corey, 2010). Is this true for cochlear bundles? In cochlear inner and outer hair cells (IHCs and OHCs), the mature complement of horizontal top connectors is established by postnatal day 12 (P12); they extend between adjacent stereocilia of both rows and columns (Goodyear et al, 2005). We therefore expect that bundle cohesion should be maintained in both directions. However in neonatal (P4) bundles, lateral coupling of OHC stereocilia appears low (Langer, Fink et al., 2001). Does this improve with development or are cochlear bundles fundamentally different from frog?

Methods
We cultured organs of Corti from P4 to P12 gerbils. A stiff probe with a blunt, ~0.5 μm tip was used to push IHC and OHC stereocilia in the excitatory direction. Stimuli were slow sinusoids of 120, 240, and 480 nm, corresponding (depending on bundle height) to 1-14 degrees. Stimuli were delivered at multiple locations along the tallest stereociliary row, and along the stereociliary height. Displacements of individual stereocilia were recorded with a CCD camera and analyzed with cross-correlation methods (Karavitaki and Corey, 2010).

Results
With data from 111 cells, we found that lateral coupling among stereocilia of the tallest row is weak. Movement in the excitatory direction propagated laterally by only a few stereocilia and the bundle did not move coherently. This was true for IHC and all three rows of OHC stereocilia, for all developmental ages, all cochlear turns and all stimulus positions and amplitudes tested.

Conclusion
The lateral coupling within the stereocilia of the tallest row is weak, suggesting that the function and molecular composition of horizontal top connectors are different from those in bullfrog hair cells. This also raises concern for current stimulus methods, which involve glass probes that are often small compared to the bundle width. Our data suggest that only stereocilia in contact with the probe will be stimulated, and delivery of the stimulus to the remaining stereocilia will be weak and therefore not homogeneous. To mimic the simultaneous and equal OHC stimulus delivered by the overlying tectorial membrane to stereocilia of the tallest row in vivo, better stimulation methods must be developed.

Electrical coupling of mouse adult inner hair cells measured in situ
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Background
Inner hair cells (IHCs) of the mammalian cochlea use ribbon synapses to signal rapidly to the afferent dendrite. Although much functional information has been obtained from relatively immature rodent cochlear coils, it has proved more difficult to record from adult hair cells as isolation of the cells can be problematic. We have overcome this issue by patch-clamp recording from adult hair cells of the mouse in the isolated temporal bone. We have used fluorescent probes to investigate calcium signalling at the synapse using simultaneous imaging with a multiphoton confocal laser scanning confocal microscope (2PCLSM).

Methods
Cochlear inner and outer hair cells were visualised in the intact isolated temporal bones of mice aged P25 to 10 mo. The cochlea was opened to reveal the 10-15 kHz cochlear region with the cells in identifiable positions and orientation. The chamber was superfused with (in mM): Na, 142; Ca, 1.3; K, 4; Cl 147; 0.7 phosphate; hepes 10, to pH 7.3 at 24°C. Cells were recorded in the whole cell tight seal mode, the pipette including 0.5, 5 or 10mM EGTA and 0.25 mM OGB-5N (excited at 935 nm by the 2PCLSM beam). Intracellular cesium was used to reduce large outward currents to less than 1.5nA at 0 mV.

Results
Distinct calcium entry sites, ‘hotspots’, could be observed on depolarizing the IHC from -60 mV. With frame rates of 80 Hz, z-sectioning of the cell showed a basal pole distribution of hotspots with the largest calcium rises on the modiolar side of
the cell. The precise kinetics and amplitude of the responses depended upon the pipette buffer concentration. In the apical region it was often found that cells were both dye and electrically coupled with up to 9 cells sometimes being found filled with OGB-5N. Calcium responses in the coupled cells could be observed on pipette depolarization; the kinetics generally differed from that of the recorded cell. The electrical input capacitance scaled with the number of coupled cells. Modelling the system suggests that the coupling resistance was less than 10MΩ.

Conclusion
Adult IHCs can be coupled. Comparison of the calcium signal in groups of coupled IHCs indicates that the endogenous buffer is excluded by the coupling pores. The approach allows visualization of the calcium kinetics of an unperturbed adult ribbon synapse.

The properties of short hair cells in the avian basilar papilla
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Background
The origin of frequency selectivity in the avian basilar papilla is incompletely understood. The tall hair cells (THCs) are electrically tuned over part of the frequency range but little is known about the short hair cells (SHCs).

Methods
To address this problem, we made patch clamp recordings from SHCs in an isolated chicken basilar papilla preparation (E18 to P5; T = 33 degrees C) at fractional distances along the papilla from the apex, d = 0.1-0.6. SHCs were identified by lower surface density and location less than 10 rows from abneural edge.

Results
SHCs are analogous to mammalian outer hair cells in having a large efferent but sparse afferent innervation and are distinguished by possessing an A-type inactivating K⁺ current (Murrow 1994). We found that SHCs are also electrically tuned by a BK Ca-activated K⁺ conductance, but such tuning is optimal only at resting potentials positive to -40 mV where the A-current is inactivated. The depolarized resting potential stems from a standing inward current through the mechanotransducer (MT) channels, whose resting open probability, Pr, is about 0.35. Two factors produce the high Pr: a low endolymph calcium (0.24 mM) and high intracellular calcium buffering. With perforated patch recordings using the MT current Pr as an assay, we estimated the intracellular calcium buffer as equivalent to 0.5 mM BAPTA (d = 0.4). To confirm the high levels of buffer, the distribution of calcium binding proteins in SHCs was examined with post-embedding immunogold labeling in transmission electron micrographs. Heavy stereociliary labeling was seen with antibodies to both calbindin-D-28K and parvalbumin-3 and displayed an apex to base gradient matching the tonotopic gradient in the MT current amplitude. The stereociliary concentrations of both calcium-binding proteins were estimated by calibrating immunogold counts against gels containing defined amounts of the proteins (Hackney et al. 2003).

Conclusion
We conclude that avian SHCs resemble mammalian outer hair cells in their large calcium buffer content, high resting open probability for MT channels and depolarized resting potential. However, the dichotomy between SHCs and THCs is less marked than between mammalian inner and outer hair cells. Funded by RO1 DC01362 from NIDCD to RF

Pitch perception in cochlear implant users with square wave iterated rippled noise
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Background
Manipulation of temporal (rate) pitch mechanisms in cochlear implant (CI) users through carrier rate modulation may expand the potential number of distinct pitch percepts available beyond that which is available through place pitch coding alone. Rate pitch has the advantage of being able to provide a continuum of pitches on a single electrode up to approximately 300 Hz. However, this limit varies greatly between subjects and within subjects (for different electrodes), with a few exceptional performers being able to distinguish rate pitch cues up to at least 1000 Hz. Square wave iterated
rippled noise (SWIRN) can be created by delaying white noise and adding it back to the original signal and transmitting it to a CI user by direct electrical stimulation.

Methods
Ten CI subjects were asked to identify the higher tone of two tones presented with direct electrical stimulation using SWIRN. The base SWIRN rates ranged from 100 Hz to 500 Hz; target SWIRN rates were 35% higher. The carrier stimulation rate was 15 kHz. Using loudness-balanced stimuli, pitch ranking was measured for three base modulation rates on groups of 3 (one basal, middle and apical electrode), 6 (two basal, middle and apical electrodes) and 11 (every other electrode) for several subjects using Cochlear Nucleus CI512 and CI24R devices. The order of the electrodes and the current level on each electrode was randomized in each block.

Results
Preliminary results demonstrate little variability across subjects, frequencies and electrode groups. Subjects seem to be able to accurately rank SWIRN stimuli in all three groups of electrodes in the tested frequency range. Midpoint comparison shows that the perceived pitch increases linearly as the modulation rate is increased from 100 to 500 Hz.

Conclusion
These results suggest that sound processing strategies that utilize SWIRN-based stimuli may lead to improved pitch and music perception in CI users.

Investigating the Effect of Training with Amplitude Modulated (AM) Tones on Tone-Vocoded Speech Perception
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Background
In vocoded speech, temporal-envelopes are extracted from broad frequency regions and used to modulate tone- or band-limited noise carriers. Such a manipulation removes temporal fine-structure and degrades spectral content whilst preserving temporal-envelope cues and, if a sufficient number of frequency channels are used (usually ≥6), retaining intelligibility. We asked here whether training on a task requiring the explicit use of temporal-envelope cues but no speech content will transfer to 4-channel vocoded speech, which is unintelligible when initially heard.

Methods
Identification of vocoded vowel-consonant-vowel (VCV) stimuli was assessed before (pre-test) and after (post-test) two sessions of training on amplitude-modulation (AM) tasks. In order to reduce any interaction between experimental effects arising from AM-based training and the rapid learning that occurs when listeners are first exposed to vocoded speech, participants were trained on vocoded VCV-identification prior to the pre-test session. During the training phase, one group (N = 18) practiced an AM-detection task, a second group (N = 18) practiced an AM-rate discrimination task, and a third group (N = 13) served as untrained Control.

Results
Although there was no significant group difference, both AM-training groups (but not Controls) showed significant improvement (~4%) on vocoded VCV identification. Analysis of the confusion matrices showed differences in the specific groups of consonants indexing training-induced changes.

Conclusion
This preliminary demonstration that training on temporal-envelope cues can transfer to the identification of vocoded speech is now being investigated using a more extended AM training regimen.

Frequency Shifts Enhance Tones within Chords, in a Special Way
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Background
In a chord of equal-SPL pure tones, one of the tones can be made to pop out perceptually by presenting, before this chord, a copy of it in which the target of the pop-out effect is attenuated, thus producing a spectral notch. A subjectively
similar pop-out or "enhancement" effect is obtained when the target tone is shifted in frequency rather than in SPL. Erviti et al. (J. Acoust. Soc. Am., 2011) reported evidence that the mechanism of enhancement is not the same for these two types of change. Here, we report other findings supporting that view.

Methods
Enhancement was measured using short-duration (at most 100 ms) inharmonic chords varying randomly in frequency content across trials and a "present/absent" task assessing the audibility of their individual components (see, e.g., Erviti et al., 2011). The second of the two successive chords presented on each trial always had a flat spectral profile. Its components were either matched in SPL to the non-target components of the first chord, or 10 dB louder than them. Enhancement was elicited either by filling a spectral notch existing in the first chord ("Profile change" condition) or by a 1-semitone shift in one tone ("Frequency change" condition); for each listener, the depth of the spectral notches (in dB) was initially adjusted so as to obtain the same enhancement magnitude in these two conditions, when the non-target tones had a constant SPL within trials.

Results
The main result (see the Figure) is that changing the SPL of the non-target tones within trials disrupted enhancement to a larger extent in the "Profile change" condition than in the "Frequency change" condition. It is also noteworthy that, when the non-target tones did not change within trials, the notch depth needed to match the enhancement produced by 1-semitone frequency shifts was as large as 11.5 dB, on average. According to the excitation-pattern model of Moore et al. (J. Audio Eng. Soc., 1997), this implies that, for a given change in the excitation of auditory filters, enhancement is stronger when it is elicited by a frequency shift than when it is elicited by filling a spectral notch.

Conclusion
While the enhancement effects resulting from changes in spectral profile are usually explained in terms of neural adaptation, this does not seem to be a sufficient explanation for the enhancement effects resulting from frequency shifts.
Using Training to Overcome the Switch Cost Induced by Stimulus Randomization on Auditory Temporal-Interval Discrimination
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Background
Alternating between two simple cognitive tasks is difficult and comes at a cost (the switch cost). In auditory perception, an example of a switch cost is the significant elevation in temporal-interval discrimination thresholds when two distinct standard temporal intervals are presented randomly trial-by-trial (roving) compared to separately in different blocks of trials (consecutive). Our goal was to determine if the cost of roving could be alleviated with practice, either with the randomized stimuli or with each stimulus in isolation.

Methods
Naïve normal-hearing adults practiced temporal-interval discrimination of two base intervals (100 and 350 ms, marked with 15-ms 1-kHz or 4-kHz tones), 720 total trials per session for six sessions in either a roving (n=16) or a consecutive (n=16) regimen. In the roving regimen, the two temporal intervals were randomly presented throughout each training session for a total of 360 trials each. In the consecutive regimen, in each session, all 360 trials of one interval were completed before training on the other interval began. A control group (n=16) completed pre- and post-training sessions only. We compared learning (on the trained intervals in the trained regimen) and generalization (to the other regimen and to untrained stimuli) between the two regimens.

Results
Both regimens yielded significantly greater improvement compared with controls on at least one of the two trained intervals, and generated similar patterns of generalization to untrained stimuli. However, the amount of learning on the consecutive regimen (estimated with effect size) was greater than on the roving regimen. Most importantly, learning did not generalize across regimens. Training on the consecutive regimen did not improve roving thresholds or vice versa. Although training on the roving regimen brought roving thresholds in line with naïve consecutive thresholds, it did not lead to improvement on consecutive thresholds.

Conclusion
The performance cost of randomly switching between two auditory temporal intervals can be overcome with multi-session training with that randomization (roving), but not with training without randomization (consecutive). Further, training with randomization does not aid performance without it. Thus, it appears that presumed improvements in stimulus encoding (after consecutive training) do not reduce the cost of switching between stimuli, and that reductions in the switch cost (after roving training) do not improve stimulus encoding. Rather, these results suggest that the switch cost between stimuli is mediated by processes separate from those that affect basic stimulus encoding.
Frequency-Modulation Vowel Maps in Normal-Hearing and Hearing-Impaired Listeners

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Background

Human singing voices are not steady in pitch but typically contain coherent frequency fluctuations over time. The perception of natural voice vibrato thus relies on the presence of coherent frequency modulation (CFM). Sensorineural hearing loss is known to affect the ability to detect changes in frequency, presumably leading to degraded voice perception. This study investigated the ability of normal-hearing (NH) and hearing-impaired (HI) listeners to perceive a sung vowel by adding CFM to a steady complex tone. The aim was to determine the extent of the “vowel map” along the FM-rate and FM-excursion dimensions in NH listeners, and whether this area for FM-based singing-voice perception is affected by hearing impairment.

Methods

The ranges of CFM rates and excursions leading to voice formation were determined in a group of NH listeners and a group of HI listeners with moderate to severe sensorineural hearing loss. The listeners adjusted either the FM rate or the FM excursion applied to the synthesized vowel /oh/ in an adaptive “yes/no” procedure, while the other FM parameter remained fixed. In each presentation, the listeners were asked to judge whether the sound corresponded to a natural singing voice. The lower and upper FM-excursion thresholds were determined for fixed FM-rates between 3 and 8 Hz, and the lower and upper FM-rate thresholds for fixed FM-excursions between 21 and 91 cents.

Results

In NH listeners, adding CFM to an unmodulated complex tone was sufficient to evoke the perception of a singing voice for FM rates between 4.3 and 7.4 Hz and FM excursions between 15 and 64 cents on average. In contrast, HI listeners typically exhibited broader vowel maps than NH listeners, whereby their maps were shifted towards higher FM excursions and extended towards lower FM rates than those of NH listeners. The large across-subject variability in the HI group was not fully explained by the listeners’ audiograms or auditory-filter bandwidths at the vowel’s fundamental frequency.

Conclusion

Hearing loss was found to affect the formation of a sung vowel based on FM-rate and FM-excursion cues. It remains unclear to what extent this is attributable to deficits in FM detection or discrimination, reduced frequency selectivity, or difficulties in following the rate of frequency changes. The vowel maps determined in NH listeners may provide reference values when constructing synthetic-vowel stimuli with realistic sung vibrato.
The Ability Of Hearing-Impaired Listeners To Use Temporal-Envelope Cues Recovered From Speech Frequency Modulation

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Background

Recent studies suggest that normal-hearing listeners maintain robust speech intelligibility despite severe degradations of amplitude-modulation (AM) cues, by using temporal-envelope information recovered from broadband frequency-modulation (FM) speech cues at the output of cochlear filters. This psychophysical study aimed to assess whether cochlear damage alters this capacity to reconstruct temporal-envelope information from FM.

Methods

This was achieved by measuring the ability of normal-hearing listeners and listeners with mild-to-moderate hearing loss to identify nonsense syllables processed to degrade AM cues while leaving FM cues intact within three broad frequency bands.

Results

Hearing-impaired listeners showed significantly poorer identification scores than normal-hearing listeners. However, the deficit shown by hearing-impaired listeners was relatively modest. Overall, hearing-impaired data and the results of simulation studies were consistent with a poorer-than-normal ability to reconstruct temporal-envelope information resulting from a broadening of cochlear filters by a factor ranging from 2 to 4.

Conclusion

These results indicate that temporal-envelope reconstruction from broadband FM is an important, early auditory mechanism contributing to the robust perception of speech sounds in degraded listening conditions. These results also suggest that most people suffering from mild to moderate cochlear hearing loss can make efficient use of reconstructed envelope cues despite degradations in frequency selectivity. Still, these results suggest that poorer-than-normal frequency selectivity impairs somewhat temporal-envelope reconstruction mechanisms.
Hearing speech in noise from a single source before and after surgical improvement of congenital conductive hearing loss
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Background
Patients with congenital aural atresia are born with a conductive hearing loss due to malformations of the external and middle ears. Surgery can improve hearing in the atretic ear to within normal limits. Atresia is thus an opportunity to evaluate roles of early conductive hearing loss on binaural processing. Here we investigate post-operative emergence of summation (binaural redundancy), the advantage of hearing with two ears rather than one when an identical signal arrives at both ears simultaneously. This simple benefit has been estimated at about 3 dB in normal listeners.

Methods
We tested 12 patients between the ages of 6 and 53 years with an average 64 dB (± 14 dB sd) unilateral conductive hearing loss, normal hearing in the non-atretic ear, and normal bone conduction in the atretic ear. Surgery improved pure-tone averages by 30 dB to post-operative PTA of 34 (± 10) dB. Before surgery and 5 weeks later, single speaker HINT tests evaluated speech reception in quiet and noise. Patients faced one central speaker. Levels of sentences were adaptively varied to determine a reception threshold. This was done first in the quiet and then with simultaneous multi-talker babble at 65 dB from the same speaker.

Results
After surgery, thresholds for speech in quiet improved 3.4 dB, and thresholds in noise improved 0.8 dB, both statistically insignificant (1-tailed p values of .1 and .2 respectively). Patients performed worse than normal in quiet and in noise (1-tailed p values of .001 and .037 respectively). There was a significant effect of age on performance in noise ($r^2$=.58) but not in quiet ($r^2$=.01). Pooling pre- and post-op results in quiet across ages, the patients were 10 (±2.2) dB worse than normal. Converting results to dB-worse-than-normal, atresia patients did better in noise (relative to normal) than in quiet (p=.008).

Conclusion
We see the expected summation effect (3 dB) in quiet but not in noise. Better performance in noise than quiet (relative to normal) might arise because these patients likely developed skills attending to threshold-level speech in noise without any binaural benefit. Since both signal and noise came from the same speaker, we don’t understand why performance in quiet remains worse than normal (by an effect size > 4). A bold speculation might be that having two ears somehow helps you learn to hear out of one ear.
Behavioral Discrimination of Ultrasonic Vocalizations in CBA/CaJ Mice

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Background
Previous studies have attempted to categorize ultrasonic vocalizations (USVs) from mice using spectrographic analysis and human-based classification (e.g. Grimsley et al., 2011; Portfors, 2007). However, relatively little research has focused on the perception of these different call types. Results from Holmstrom et al. (2010) illustrate that neurons in the inferior colliculus respond differently to an array of USV categories, suggesting that the auditory system uses specific patterns for encoding different call types. However, until now, no study has examined how well different USVs are discriminated by awake, behaving mice.

Methods
Both the perceptual saliency of USV categories and the impact of different call manipulations were measured here. CBA/CaJ mice were trained to discriminate target USVs from a different USV presented in a repeating background. The repeating background signal in a session was one of five different call types that were previously used by Holmstrom et al., 2010: two frequency modulated up-sweeps, one produced by a male and the other by a female, as well as harmonic jump calls that varied in the number (0-2) of jumps found within the call. The discrimination targets in a session included the four non-background USVs and spectrotemporal manipulations of the background call. The spectrotemporal manipulations were removing the frequency modulation, shifting the frequency up and down ten and twenty percent, shortening and lengthening the call by a factor of two, and reversing the signal.

Results
Interestingly, many calls within human-created ‘categories’ were easily discriminated from one another while the mice were unable to discriminate other between-category calls, suggesting that current conceptions of USV categories may not be accurate for awake, behaving mice. The discrimination performance on the spectrotemporal manipulations indicates that some call parameters are more important than others for USV perception. For most calls, large frequency shifts (+/- 20%) resulted in high discrimination performance, suggesting that frequency information may be important for ultrasonic communication. Manipulating the duration of the USV appears to have disparate impacts among harmonic jump calls and up-sweep USVs, suggesting that the role of temporal cues may differ between call types. Surprisingly, reversing the call and removing the frequency modulation had a substantial reduction in discrimination performance for most USVs.

Conclusion
These results suggest that frequency modulation and contour have little impact on how USVs are perceived, and imply that categories emphasizing these differences (e.g. upsweeps vs. downsweeps) are not relevant for USV perception.
The Pitch of the Broadband Temporal Envelope: Effects of Modulation Rate and Shape

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Background
Pitch perception of an amplitude modulated (AM) signal depends on the modulation rate, the carrier bandwidth and the modulation shape. While much attention has been paid to the effects of rate and bandwidth of the carrier, fewer studies have investigated effects of modulation shape. Moreover, cochlear implants (CI) users rely primarily on temporal envelope cues for complex pitch perception. Attempts to improve coding of the fundamental frequency (F0) for CI listeners have used peakier modulation envelopes. In the present study we examined the temporal pitch percept conveyed by such envelopes in normally hearing (NH) listeners.

Methods
We investigated NH listeners’ sensitivity to rate and shape of the broadband temporal envelope of noise stimuli. The stimuli were either sinusoidally amplitude modulated (SAM) noise or noise that received the broadband temporal envelope of sine-phase harmonic complexes based on a fundamental frequency equal to the rate of SAM. These harmonic envelope modulations (HEM) varied in peakiness depending on the number of partials in the complex modulator.

Results
Experiment 1 showed that both shape and rate had some influence on loudness, which could be released by a level roving ranging over 3 dB. Experiment 2 measured discrimination of the shape of noise stimuli modulated at the same rate. Sensitivity worsened at higher rate and at higher peak factor. Experiments 3 and 4 showed that even though rate dominated the sensation of temporal pitch, shape had an influence: peaky modulations tended to sound lower in pitch than sinusoidal ones. Experiments 5 and 6 measured listeners’ sensitivity to discriminate modulation rates keeping a constant shape. Difference limens were lower for the HEM than for the SAM stimuli at both 100 and 200 Hz, but remained much larger than those obtained with full harmonic complexes.

Conclusion
First, the sensitivity of NH listeners to temporal pitch can be improved by using modulations peakier than sinusoidal. Second, temporal pitch is not entirely governed by modulation rate: modulations peakier than sinusoidal tend to sound lower in pitch. Third, NH listeners are able to discriminate temporal envelopes that are modulated at the same rate with different shapes. Thus, CI users might well use both rate and shape of modulation to perform a pitch discrimination task. A larger focus on modulation shape might help to improve F0-coding in CIs.

Phase Effects in Speech Recognition Masked by Harmonic Complexes

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Background
Compression by the basilar membrane (BM) helps to listen in extremely short temporal dips and facilitates tone detection in masker waveforms that are deeply modulated. Its role for speech understanding in a background of other voices remains however unclear.

Methods
Experiment 1 measured speech reception thresholds (SRTs) for an unprocessed male and a female voice masked by broadband or speech-shaped harmonic complexes with partials either in sine phase (SP) or in random phase (RP). The masker’s fundamental frequency (F0) was 50, 100 or 200 Hz. Experiment 2 measured masked thresholds (MTs) for speech-shaped noise that was restricted to the regions of resolved or unresolved partials of the same maskers used in experiment 1.

Results
In experiment 1, SRTs were lower for SP than for RP maskers only at 50-Hz F0. This masking release (MR) was absent at 100-Hz F0 and was negative at 200-Hz F0. The results were similar whether the maskers were broadband or speech-shaped and whether the target talker was male or female. In experiment 2, MTs were similar for SP and RP maskers in the region of resolved partials, and lower for SP than RP maskers in the region of unresolved partials at both 50- and 100-Hz F0. It appears that the MR observed at 100-Hz F0 did not transfer from MT to SRT because the SRT was lower than the range of target-to-masker ratios where compression is potentially beneficial. In other words, speech can often be understood at 50% intelligibility without the information located in unresolved regions of the masker. Compression can
play a critical role but only at supra-threshold performance. Psychometric functions were reconstructed from the adaptive tracks of experiment 1 to extract thresholds at 65% and 80% intelligibility. Only at 80% intelligibility, was a MR observed at 100-Hz F0.

Conclusion
Overall, these results suggest that BM compression contributes weakly to listening in extremely short temporal dips present in steady-state speech-shaped periodic maskers at 65 dB SPL.

Interactions of Pitch and Timbre: How Changes in One Dimension Affect Discrimination of the Other
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Background
Variations in spectral shape, perceived as timbre changes, can lead to poorer fundamental frequency (F0) or pitch discrimination. Less is known about the effects of F0 variations on the discrimination of spectral shape. The present study used bandpass-filtered complex tones to examine both the effect of changes in spectral centroid on pitch discrimination, and the effect of F0 changes on timbre discrimination. Variations in the two dimensions were equated for salience by normalizing changes relative to individual subjects’ difference limens (DLs). The two aims of the study were 1) to determine whether the interactions between pitch and timbre were symmetric, and 2) to assess the effects of musical training on listeners’ ability to ignore variations in irrelevant perceptual dimensions.

Methods
The thirty subjects were divided into two equal groups of musicians and non-musicians. In Experiment 1, the subjects’ task was to compare sequentially presented tone pairs that differed in either pitch or timbre and to judge which was higher. The DLs obtained for both tasks were used in subsequent experiments. In Experiment 2, F0DLs were measured as a function of the size of random variations in spectral centroid, and vice versa. In Experiment 3, sensitivity was measured as the target parameter and the interfering parameter varied by the same amount, in terms of individual DLs.

Results
Both pitch and timbre DLs were affected by random variations in the non-target dimension. The amount of interference observed was similar for both pitch and timbre dimensions. Although musicians had lower (better) F0DLs than non-musicians on average, the amount of interference produced by random spectral variations was similar for the two groups. In addition, there was no significant difference in spectral centroid (timbre) DLs between musicians and non-musicians with or without random variations in pitch. Overall, performance was better when the random non-target variation was in the same direction as the target variation (e.g., when an upward movement in timbre was paired with an upward movement in pitch).

Conclusion
Difference limens for both pitch and timbre are strongly, and similarly, affected by random variations in the non-target dimension. Although musical training does not seem to reduce this interference. The results confirm that pitch and timbre are not easily separable as perceptual dimensions of hearing, and that directional changes can be confused across the two dimensions.

Masker Profiling Yields Insights into Mechanisms of Masking
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Background
Signal detection in noise is assumed to be a function of signal-to-noise ratio in the frequency region of the signal, but there is little consensus regarding specific decision strategies.

Methods
Thresholds were determined for eight normally hearing listeners for detection of a signal (2000-Hz sinusoid, 100 ms long, and 20 ms ramps) centered in an 8-ERB wide broadband noise (1219-3200 Hz, 300 ms long, 5 ms ramps) presented at
47, 62, and 77 dB SPL. An independent level rove of +/- 8 dB in 1 dB steps was then applied to individual half-ERB wide bands of the masker while the signal was fixed at either +4 or +8 dB above threshold. In each condition, listeners ran 1000 trials in a Yes-No paradigm. Perceptual weights for masker bands were obtained using a reverse correlation method, separately for signal and no-signal trials, such that a positive weight indicates a tendency to vote “Yes” when the level of the masker band is higher. In a new masker profiling analysis based on the overall framework of signal detection theory, trial-by-trial data were divided into four categories of responses namely, Hits, Misses, False Alarms, and Correct Rejections and the mean level of each masker band within each response category was calculated.

Results
Perceptual weighting patterns reflect auditory filter shapes and indicate that listeners respond “Yes” on signal trials when the level of masker bands near the signal is low, but respond “Yes” on no-signal trials when the level of those bands is high. The four-category masker profiling analysis provides a basis for the interpretation of perceptual weights by showing the average masker levels resulting in the “Yes” and “No” decisions that contribute to each weight.

Conclusion
Lower mean levels of masker bands near the signal in Hits and higher levels for those bands in Misses suggest that detection is based on signal-to-noise ratio. On no-signal trials, however, listeners vote “Yes” when the level of the masker in bands near the signal is higher, consistent with an energy detector. Since signal and no-signal trials are presented randomly and listeners are blind to this information a priori, it is generally assumed that listeners’ decision strategies are similar on signal and no-signal trials, but this is clearly not the case. Use of masker profiling may provide significant information to improve models of masking.
Behavioral paradigm to measure the lower limit of pitch for complex harmonic sounds in macaques.

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**Background**

Pitch is an auditory percept related to the repetition rate of sounds but with a complex relationship to the physical attributes of sounds. To study the neural correlates associated to pitch perception using non-human primate models, behavioral measurements of pitch perception in these animals are critical. In humans, the range for pitch perception is approximately 30-5000 Hz but this range is currently unknown in monkeys partly because of the difficulty to train monkeys for auditory tasks (Brosch et al., 2004). In this study, we developed and applied a behavioral paradigm to measure the lower limit of pitch for complex harmonic sounds in macaques based on changes in the discrimination threshold across the values of repetition rate (Krumbholz et al, 2000).

**Methods**

We trained three rhesus monkeys (Macaca mulatta) using a “go/no-go” procedure to release a touch bar sensor at each detection of a change in repetition rate. Stimuli were harmonic complexes (cosine and alternating phase) with repetition rates in the range of 8 to 128 Hz. The sounds were played diotically through earphones. A trial began with an initiation interval during which a trial could be initiated by contacting the touch bar. Upon contact, a holding period (1-3 sec) began. During this period, the standard harmonic complex sound was played and followed by a stimulus change interval during which a comparison sound with a higher rate was presented. The monkeys responded with bar release to detected changes in rate (test trials) and were rewarded for hits and for correct rejections (holding the bar during catch trials). We used an adaptive staircase procedure with 3 trials (2 test trials and 1 catch trial) per step (Moore and Fallah, 2004) to determine the thresholds.

**Results**

Our paradigm was used to estimate the lower limit of pitch in the 3 animals who successfully learnt to detect a change in repetition rates. We found that each animal shows a lower threshold (<5%) for higher rates (> = 32 Hz).

**Conclusion**

The lower limit of pitch in rhesus monkeys (~30Hz) seems very similar to that of humans. This work allows the behavioral definition of a pitch threshold against which BOLD responses to pitch-associated stimuli at different rates can be compared (Baumann et al., 2012): a predicted property of a neural correlate of pitch is an existence region only above lower limit.

Effects of noise reduction on AM discrimination for hearing-impaired listeners

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**Background**

Noise-reduction (NR) algorithms are employed in digital hearing-aid devices to improve the listening experience of the user in noisy backgrounds. Although these algorithms may increase the signal-to-noise ratio (SNR) in an ideal case, they generally fail to improve speech intelligibility. However, due to the complex nature of speech, it is difficult to disentangle the numerous effects of noise reduction which may underlie the lack of speech benefits.

Ives et al (2012) examined the effect of a NR algorithm on the ability of normally-hearing (NH) listeners to discriminate a basic acoustic feature known to be crucial for speech identification, namely amplitude modulation (AM). They found that NR slightly improved discrimination at higher SNRs. The goal of the present study was to assess whether the benefit of NR on AM discrimination was present for hearing-impaired (HI) listeners.

**Methods**

The discrimination of complex AM patterns was measured for 10 HI listeners and 10 NH listeners using a same-different discrimination task. The stimuli were generated by modulating a pure-tone carrier by a two-component AM modulator with modulation rates centered around 3 Hz. The carrier tone was either 500 Hz or 2 kHz and fixed within a block.
Discrimination was measured for both groups of listeners (NH and HI) at 500 Hz and 2 kHz in the presence of a band-pass filtered Gaussian white noise at an SNR of 12dB. Stimuli were left as such or processed via a NR algorithm based on the spectral subtraction method. The HI listeners had normal hearing (≤20dB HL) at 500 Hz and moderate-severe hearing loss (≥40dB HL) at 2 kHz.

Results
NR was found to: (i) improve AM discrimination at both 500 Hz and 2 kHz for the NH listeners; (ii) improve AM discrimination at 500 Hz for HI listeners; (iii) have no effect on AM discrimination at 2 kHz for HI listeners. The stimuli were passed through a computational model of the peripheral auditory system. The simulation results suggest that the lack of benefit of NR for the HI listeners at 2 kHz may arise from the combined effects of higher absolute thresholds and reduced cochlear compression. Auditory filter width did not affect performance.

Conclusion
HI listeners do not benefit from NR for AM discrimination. The results suggest that this lack of benefit may arise from a poor matching between the compression stage and NR algorithm in hearing aids.

Effects of Noise on Detection of Vowels by Nonhuman Primates
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Background
Vowel detection in noisy conditions can help understand detection of general complex signals. In macaques, the thresholds for pure tone detection increase by 1 dB per dB of broadband noise added, maintaining the signal to noise ratio. In humans, vowel detection has been shown to depend on the signal to noise ratio of the most intense formant peak, but it is not clear that these mechanisms are in play in macaques which attach no semantic significance to and have no previous experience with vowels.

Methods
To investigate some of the underlying principles of detection of complex sounds, we examined detection of vowels in various kinds of noise. Three macaques (Macaca mulatta) were trained in a reaction time lever release task with interleaved catch trials to report detection of vowels presented alone and in continuous noise. We manipulated the bandwidth and the frequency range of the noise in order to evaluate the signals and noise components that influence detection. Signal detection theoretic analysis was used to determine behavioral accuracy from hit rates and false alarm rates.

Results
We studied the effect of noise on reaction times as well as detection thresholds, but found that noise only influenced thresholds. Detection of vowels shows the classic sigmoid relationship with sound pressure level. Broadband noise shifted the thresholds of vowel detection by +1 dB for 1 dB of noise. The pattern of incremental threshold shift did not change significantly when a mixture of vowels was used as background. When the bandwidth of the noise was restricted to 4000Hz, the frequency range of the vowel signal itself, it still caused a shift rate of 1 dB/dB, not significantly different from broadband noise. When the 4000 Hz noise band was translated in frequency, the shift rate decreased as the overlap between the vowel and the noise band decreased. When the noise components corresponding to the formant peaks were individually notched, no single formant was found to have predominance in vowel detection. When the bandwidth of noise was increased, the effect increased continuously but saturated beyond the second formant; increasing bandwidth while fixing the high frequency end caused small changes until both first and second formants were encompassed by the noise.

Conclusion
These results suggest that detection of vowel signals may relate to the total pressure over a partial band of the signal rather than the individual formant peak or the total pressure over the whole signal band. (NIH_R01_DC_11092)
Perception and neural representation of tones in conditions of masking release
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Background
The audibility of sounds in natural acoustic environments is often hampered due to the presence of other masking sounds. The amount of masking can be reduced due to the presence of beneficial signal properties. Psychoacoustical experiments showed that a masking release can be found in the presence of coherent intensity fluctuations across frequency (comodulation masking release, CMR) or interaural signal phase disparities (binaural masking level difference, BMLD) compared to a condition where those properties are absent. It was shown (Epp et al., ARO 2012) that a release from masking is reflected in auditory evoked potentials evaluated at constant stimulus levels and that the P2 amplitude reflects the level above masked threshold better than the physical level of the stimulus. It is hypothesized that a psychoacoustical measure of the audibility of the signal also correlates with the auditory evoked potentials when evaluated at the same level above masked threshold.

Methods
To assess the psychoacoustical measure of audibility, thresholds were measured for masked tones. Masked threshold was varied with the introduction of comodulation, interaural signal phase disparity or a combination of both. Based on the individual data, the listeners were asked to rate the audibility of a masked tone at individually adjusted physical levels corresponding to constant levels above threshold. To assess the neural activity, auditory evoked-potentials were measured for the same stimuli as used in the rating experiment within the same listeners.

Results
Psychoacoustical results indicate a growth of audibility that is increasing with increased level above masked threshold. Conditions without a release from masking show a tendency for higher audibility than conditions with a masking release. Auditory evoked potentials show an increase in amplitude with increased level above masked threshold. The analysis of the evoked potential shows differences in sensitivity between N1 and P2 to comodulation and interaural signal phase. The electrophysiological data with constant level above masked threshold are in agreement with data of Epp et al. (ARO 2012) with constant physical stimulus levels.

Conclusion
The psychoacoustical data show that audibility is mainly determined by the level above masked threshold rather than the physical level of the stimulus. The higher audibility for conditions with a masking release might be some residual influence of the overall level of the stimulus on the audibility rating. The electrophysiological data support the hypothesis that the P2 component of auditory evoked potentials correlates closely with levels above masked threshold rather than with physical stimulus levels.
Frequency Difference Limens and Cue Trading in CBA/CaJ Mice
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Background
Although much is known about the genetic makeup and physiology of the laboratory mouse, far less is known about mouse auditory behavior. Using behavioral approaches, previous researchers have examined the frequency and intensity discrimination abilities of the NMRI mouse, feral house mice, and other strains of mice with known hearing impairments. The present experiments examine the hearing of the CBA/CaJ mouse strain using operant conditioning procedures.

Methods
Seven adult CBA/CaJ mice were used in these experiments. The test stimuli used in the frequency difference limens (FDL) experiment were pure tones at 12, 16, 24, and 42 kHz presented at both 10 and 30 dB (SL). The mice were trained to discriminate a comparison tone (target) from a reference tone (repeating background). The target tones were higher in frequency than the reference stimulus. The cue trading experiment used an identification paradigm where the mice had to place tones into one of two categories. For example, if a mouse heard a 70 kHz tone with a 25 ms duration it had to respond to the left response hole, and if it heard a 30 kHz tone with a 120 ms duration, it had to respond to the right response hole. Once the mouse was able to identify these two stimulus categories accurately, a 70 kHz tone with a 120 ms duration was presented. If the mouse was using frequency to identify the tones then it should choose the left hole; however, if duration was the more important cue, then it should choose the right hole. Using this paradigm it was possible to determine whether frequency or duration was the more salient cue for mice when identifying tones.

Results
In the FDL experiment, the mice had a mean just-noticeable-difference (JND) of 3.5% Weber fraction across all four frequencies and sound levels. In the cue trading task, the mice overwhelmingly identified tones based on their frequencies rather than their durations.

Conclusion
Results from the FDL test showed that CBA/CaJ mice had similar Weber fractions as wild mice and guinea pigs but had much poorer frequency resolution compared to humans and cats. Despite this poorer resolving power, the mice used frequency cues over duration information in the cue trading task, suggesting that frequency might be the dominant cue that mice use to discriminate among various mouse vocalizations.

Matching of the Dominant Pitch of Scale Alternating Wavelet Sequences against Complex Tones with Odd Harmonics Attenuation: Decision Statistics on the Stabilized Auditory Images
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Background
Scale alternating wavelet sequences (SAWSs) has been introduced where a certain wavelet and its scaled version alternate every other cycle of the overlap-and-adding. It has been reported that the pitch of the SAWS tended to shift downwards by an octave when the differences in scaling became large. The observed pitch shift could be matched to the harmonic complex tones whose odd harmonics were attenuated. The purpose of the current study was to inquire a better decision statistics to predict the observed pitch shift.

Methods
The SAWSs were generated based on the wavelet of the impulse responses of three Japanese vowels. The original wavelet and its scaled version were overlap-and-added alternately at an cycle of 8 ms. The scaling factor ranged from 0.50 to 2.00. Odd harmonics attenuated complexes (OHAC) were used as comparison stimuli. The OHACs were harmonic complexes with a 16 ms fundamental period and with each vowel’s spectrum envelope. When the attenuation level is infinite, the OHAC becomes a harmonic complex whose fundamental frequency raises by an octave. The listeners’ task was to adjust the attenuation level until the perceived pitch became equivalent to that of the SAWS.

Results
The matched attenuation level varied systematically with the divergence in the scaling factor. Tsuzaki et al. (2012) demonstrated that the ratios between two major peaks in the summary stabilized auditory images (SAIs) were good indicators to explain the dominating pitch, while Dinther et al. (2006) reported that the difference between the area under the two dominant peaks worked well to explain the pitch saliency for the OHAC. Two scatter plots using the two decision
statistics can be obtained where the points on the plane corresponds pairs of the SAWSs and their matched OHACs. If the prediction were complete, all the points should be on the line whose slope is a unit. The RMS errors from this line were 0.269, and 0.299, respectively for Tsuzaki’s, and for Dinther’s decision statistics. This suggests that the peak ratios could be a better indicator for the salience of the dominant pitch.

**Conclusion**

Based on the results of the pitch matching task between the SAWSs and OHACs, the simulations using the SAIs demonstrated that the ratio between the height of two peaks in the summary SAI could provide a better prediction than the difference between the area under the two peaks.
On the nature of rhythm and memory for time
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Background
Internal clock models ascribe a working memory component which serves as an intermediate module between the accumulator and the comparator (Church, 1984). Previous work on temporal memory was limited and did not consider sequences with different rhythmic structures and memory loads as occurs in natural stimuli like speech and music.

Methods
Here, we developed a novel paradigm to analyze working memory for single time intervals in the context of sequences with different: (1) rhythmic structures: jitter ranging from 5-50%, (2) memory loads: 1-4 intervals, and, (3) inter-onset intervals – sub-second (500-600ms) and supra-second (1000-1200ms).

Participants listened to intervals in a sequence of clicks and were probed on their memory for a random interval after a variable delay. Response was initiated by a programmed click and participants matched the duration of the probed interval by terminating the response interval with a button press. Feedback, defined as the difference between the duration of the probed and the reproduced interval was provided.

Time matching responses were analyzed in terms of precision, or the inverse of the standard distribution of the error responses. Consistent with distributed resource models of working memory which posit that memory can be dynamically shared between one or more items (Bays and Husain, 2008), precision provides a measure of the variability of the memory for an item around its true value.

Results
Results from experiment 1 indicate that the decay of precision with decreasing temporal regularity can be best modelled as a quadratic function (n=10) and showed a significant main effect of jitter (p=0.011). Experiment 2 showed a significant effect of working memory load (p < 0.05) and a linear decay of precision with memory load (n=9). Experiment 3 did not show a main effect of jitter (p = 0.65) suggesting that sensitivity to temporal regularity is lost at supra-second intervals and also revealed a linear decay of precision with temporal regularity (n=10).

Conclusion
These results suggest for the first time that the memory for time intervals between auditory stimuli depends crucially on the context and varies with temporal structure and working memory demands. Results from these and ongoing experiments will be discussed in terms of mechanisms of timing that consider the rhythmic structure of time intervals (Grube et al., 2010; Teki et al., 2011), models of time perception (Meck, 2005; Teki et al., 2012), as well as working memory.
The role of auditory feedback on vocal pitch production in cochlear implant users

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Background
Cochlear implantation (CI) has been highly successful in improving speech perception and production in people with profoundly hearing loss. However, how CI users perceive pitch cues in speech and how these cues affect speech production remains poorly understood. In this study, we investigated two questions. First, how does vocal pitch production in CI users differ from that of normal hearing (NH) listeners? Second, how does CI-mediated auditory feedback influence vocal pitch production in CI users?

Methods
We recorded the vocal production of consonants (in the form of a+C+a words) and vowels (in the form of h+V+d and s+V+t words), the vocal pitch simulation of the word “Ma” in the pitches of seven musical notes, as well as melody singing in postlingually-deafened CI users and NH listeners. We evaluated the vocal pitch of CI users by asking NH listeners judging the pitch contour that CI users produced and related the results to their pitch contour perception data. To assess the influence of auditory feedback through the CI on vocal pitch production, we compared speech production and singing in two conditions in CI users, CI on or CI off.

Results
Results show that pitch production by CI users is significantly less accurate than by NH controls. CI users typically produced lower pitch with greater variations in comparison to NH controls. However, the performance of pitch contour produced by CI users as judged by NH listeners is relatively accurate and is significantly correlated to their pitch contour perception. Vocal pitch production with CI off was generally less stable than when the CI was on. When the CI was turned off, CI user’s vocal production was typically louder than when the CI was on. In addition, the fundamental frequency and harmonics of vowels shifted towards higher frequencies in CI off comparing with CI on condition in most CI subjects.

Conclusion
These findings show that vocal production of CI users captures the contour but not the absolute pitch of the target sounds, indicating that vocal production of CI users is affected by the impoverished auditory feedback received and the poor pitch perceived through their CIs. Furthermore, these findings suggest that optimizing auditory feedback in CI users should improve both vocal pitch production and speech intelligibility.
A Model of Bi-Modal Localization in Young Children
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Background
The Duplex theory of localization has long been the main explanation of localization. However, given the varying developmental rates of different sensory modalities in humans, it is possible that factors such as task comprehension, vision, and attention contribute to auditory localization capabilities in young children. Visual development is complete before the end of the first year of life but auditory development is more complex anatomically and continues to develop up to and beyond age twelve. Spatial attention in children is stimulus-directed up until approximately twelve years of age. Beyond twelve years old, spatial attention becomes more goal-directed. A model for the study of visual and auditory development influences on localization was designed. Localization acuity was then tested in these two sensory modalities to separate non-auditory attention and task comprehension factors from the developmental central auditory processing factors affecting sound localization in young children.

Methods
Localization testing was conducted using visual and auditory stimuli with a group of 12 adults and 3 groups of young children aged 3-, 4- and 5-years old respectively. Testing occurred in a sound-treated booth containing a semicircular array of 15 loudspeakers. Each loudspeaker had a tiny light bulb and a picture fastened underneath. Seven of the loudspeakers were used to randomly test sound and light source identification. The sound stimulus was the word “baseball”. The light stimulus was a flashing of the light bulb triggered by the digital signal of the word “baseball.” Each participant was asked to indicate the location of test stimuli upon presentation.

Results
Localization acuity was significantly better for light than sound stimuli for children and adults. When compared to adults, children showed significantly greater localization error for both sensory modalities. 3-year-old children had significantly greater sound localization error compared to that of 4- and 5-year-olds. Adults performed better on the sound localization task when the light localization task occurred first.

Conclusion
Young children can understand and attend to localization tasks. When the central processing burden is lessened, children and adults show improved localization acuity to visual stimuli over auditory stimuli. Children show poorer localization acuity than adults across sensory modalities, which may reflect differences in both attention and central processes in young children. Young children, like adults, do not rely exclusively on level cues to localize sounds. Adults can infer the location of sound sources based on prior experience localizing light sources.

How do cognitive factors interact with speech-in-noise segregation in normal hearing children and adults?
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Background
In noisy environments, it is difficult to attend to a talker while simultaneously ignoring background speech and noise. Compared to adults, children have more difficulty extracting target speech from interfering noise, and demonstrate greater variability in performance on source segregation tasks. Little is known about auditory or non-auditory factors accounting for this variability. This study tested the following novel hypotheses: semantic content of target speech influences speech intelligibility for children; measures of cognitive assessment predict performance on speech-in-noise tasks; and cognitive measures are inversely correlated with informational masking.

Methods
Four groups of normal hearing (NH) listeners were tested: Elementary school (7-10yrs), Middle school (11-14yrs), High school (15-17yrs) and Post-Secondary (18-22yrs). The auditory task consists of subjects repeating target sentences that are either semantically coherent or anomalous. Target speech is presented in quiet and with interfering speech or noise at four signal-to-noise ratios (SNRs) (-16, -8, 0, 8dB), representing the target level relative to the interferer. The target and interferers are either collocated or spatially separated. Cognitive tests assess working memory (WM) and attention, and are administered through computer-interactive methods. WM is assessed using a List Sorting Test requiring recall and sequencing. Attention is assessed using the Flanker Test, involving focusing on one stimulus while inhibiting another, and
the Dimensional Change Card Sort Test, measuring flexibility by matching pictures according to various dimensions (e.g. color, shape).

**Results**
Preliminary results indicate the percentage of words correctly identified at each SNR, in each spatial configuration, is greater for target sentences that are semantically coherent than those that are anomalous. Percent correct decreases at lower SNRs, especially when both the target and interferers are speech and sources are collocated. Data from cognitive measures will be reported to determine whether the hypotheses are confirmed.

**Conclusion**
This is the first study to evaluate how children's WM and attention predict individual differences in ability to hear speech in noisy environments. Consistent with adult data, semantically coherent sentences are more accurately reported, indicating semantic content influences intelligibility of target speech. Performance on these tests improves with age, suggesting they might provide a useful tool for tracking the developmental trajectory of source segregation in NH listeners. The tests used in this study may be informative in assessing individuals with hearing impairment and who use assistive listening devices.

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**Hearing the elephant in the scene: A computational model of bottom-up auditory attention for auditory scene analysis**

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**Background**
As we listen to our environment in our daily lives, our brains are constantly extracting information from the incoming sound waveforms. One aspect of this mechanism is the detection of sounds that appear to be outliers within the context of their surrounding sounds. We hypothesize that as the sound statistics change over time, variance from these statistics drive bottom-up attention processes that direct our focus to certain events in a scene.

**Methods**
Here, we propose a model of bottom-up attention around a predictive coding framework. We use a rich high dimensional feature space that aims to capture the various characteristics of sounds that can steal the spotlight of attention. Among each feature axis, the statistics of the signal is learned; leading to a prediction for the next time instance. We obtain an estimate of saliency across time for every feature as a result of how well the real signal matches the predictions. The streams are fed to an interaction mechanism which brings the information across different features together in accordance with human experiment results. The result is a prediction of the times when an event occurs that is likely to grab the attention of a listening person.

**Results**
We present both human experiment results and computational model results using the same set of stimuli, and discuss the strengths and weaknesses of the attention model in matching the human experiment results.

**Conclusion**
We have presented a computational model of bottom-up auditory attention, introducing a predictive coding framework for the first time. Our model provides an efficient way to computationally find deviants in sound signals, and lays the foundation for investigating more sophisticated aspects of auditory attention.
Psychophysiological analyses of the effects of phase information on perceptual outcomes for normal-hearing listeners
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Background
Previous studies have demonstrated the importance of encoding phase cues for speech perception, and it is generally agreed that deficits in encoding phase cues result in degraded speech perception, particularly in noise. However, it is still not clear to what extent disrupted phase encoding is related to a reduction in processing of complex stimuli. The goal of this study was to understand the systematic relationship between the degree of accurate phase information available for listeners and perceptual outcomes. The encoding of such phase cues in the auditory nerve was investigated using a computational auditory model.

Methods
A phase randomization vocoder was used to systematically vary the degree of phase information available for listeners from 0 to 100%. To determine the effect of phase randomization upon speech perception, sentence recognition was measured for normal-hearing listeners in the presence of a steady or modulated masker. To further understand the nature of the effect of phase randomization, psychoacoustic performance was also evaluated. Spectral-ripple discrimination was used to evaluate spectral processing and Schroeder-phase discrimination was used to evaluate temporal processing as a function of the degree of phase randomization. Neural cross-correlation coefficients were computed to quantify neural envelope and temporal fine structure (TFS) coding in model auditory nerve spike trains.

Results
The phase randomization vocoder successfully varied the degree of phase cues available in the signal as well as in the auditory nerve. An almost linear relationship was observed between the broadband TFS of original and vocoded speech signals as a function of the degree of phase randomization. Results from the computational auditory model showed that neural TFS coding was systematically degraded as the phase was disrupted in the acoustic speech signals. A change in neural envelope coding was also observed, but these changes in neural envelope and TFS coding were independent from the effect of the number of vocoder bands. The perceptual outcomes utilizing the randomized phase information were shown to be different for different tasks. In general, performance improved as more accurate phase information was provided, but the pattern of improvement was dependent on testing conditions (masker type or number of bands).

Conclusion
The present study demonstrated that disrupted phase in the acoustic signal leads to the disrupted representation of neural coding of those cues, and this disrupted neural coding results in a reduction in perceptual performance.

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reduced audibility and poorer frequency resolution were achieved through attenuating and spectrally smearing the stimulus, respectively.

Results
Listeners with either hearing impairment or simulated impairment experienced comparable benefit of dichotic presentation as normal-hearing listeners without a simulated loss. Asynchronous glimpsing was evident in all listening groups, indicating an ability to integrate dichotic speech information across time, frequency, and the two ears.

Conclusion
By varying levels of spectral smearing, the present experiment explored the impact of impaired frequency resolution on noise-masked speech perception. Preliminary results indicate that hearing impaired listeners can benefit from dichotic presentations in an asynchronous glimpsing task, and therefore, may provide impetus for new processing strategies in bilateral auditory prostheses.

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Learning to understand degraded speech: effects of age, masker configuration, and training procedure
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Background
This work represents a first step towards developing a game-based educational curriculum for improving communication. The experiments were designed to evaluate how training with an audio-game compares to training with videogame and testing-based regimes. Speech reception thresholds (SRTs) were measured before and after training under a variety of different masker configurations. The masker configuration used during training was systematically manipulated to investigate the transfer of training.

Methods
Two experiments consisting of three stages (pre-test, training, and post-test) were conducted. During the pre- and post-tests, SRTs were measured with vocoded speech targets and vocoded (1) on-frequency speech, (2) on-frequency noise, and (3) off-frequency speech maskers that were either co-located or spatially separated from the target. In Experiment I, there were five groups. One group of younger listeners trained with a videogame. The other four groups (two older and two younger) trained with an on-frequency speech masker that was spatially separated from the target with the training being either audio-testing or audio-game based. In Experiment II, three groups trained with the audio-game and the masker configuration used during training varied amongst the groups.

Results
All groups demonstrated statistically reliable improvements in SRTs. The younger groups had a lower overall SRT, however, the pattern and amount of learning was independent of age. There was no difference in the pattern of learning between the audio-testing and audio-game groups. There was a statistically reliable difference in the pattern of learning between the videogame and audio-game groups. The overall trend was towards greater learning in the audio-game group, however, in some conditions the trend was towards greater learning in the video-game group. Further, the maximal amount of learning was not reliably observed in the trained stimulus condition and there was no obvious relationship between the pattern of learning and the training condition.

Conclusion
Audio-game based training can be just as effective as audio-testing based training. Further, the amount of learning is independent of age. This suggests that audio-game based training may present a viable means of alleviating some of the burdens of hearing impairment. The complexity of the learning patterns and their dependence on the masker configuration used during training suggests that optimizing a training-based intervention will be non-trivial. [This work was supported by the NIHR and Royal Society.]
Modeling Speech Intelligibility Performance with Spatially Separated Interferers in Normal Hearing and Hearing Impaired listeners
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Background
The Equalization-Cancellation (EC) model (Durlach 1963, JASA) has been used extensively to predict the outcomes of various binaural processing experiments. In an ongoing experimental study, speech reception thresholds (SRTs) were measured monaurally and binaurally in NH and HI listeners with monotonic speech stimuli, with a target in front and two maskers laterally displaced either with one in each hemisphere (symmetric) or both in the same hemisphere (anti-symmetric). The purpose of this work is to see how model predictions relate to the experimental data, including the effects of various model parameters.

Methods
Speech sentences were bandpass-filtered into 1/3-octave narrow bands and jittered in time and intensity. These jittered narrow bands comprise monaural channels to the model's decision stage. For each filter band, a binaural channel is created consistent with the EC model modified to process short time-sections of the stimuli separately, each with its own equalization and cancellation processing to minimize masker energy. The resulting processed time sections were combined to form a signal with an improved signal-to-noise ratio (SNR). It was assumed that the speech intelligibility index (SII) could be calculated using the channels with the best SNR in each frequency bin. In this study, the time-slice window duration and the time- and intensity-jitter standard deviations were varied to see how the threshold predictions changed.

Results
Experimentally measured binaural SRTs were variable, up to about 8 dB lower than the monaural SRTs. Symmetric binaural model predictions with normal jitter parameters varied between 1.5 and 9 dB below monaural predictions, depending on window length (between 10 and 1600 ms). Anti-symmetric binaural predictions were 6 dB below monaural. When the jitter was decreased by a factor of four, symmetric predictions were relatively insensitive, while anti-symmetric SNRs decreased by 3 dB. Increasing the jitter made the predictions equal to the monaural predictions.

Conclusion
Binaural advantage data could be fit by the model for either masker configuration. The nature of the parameters that varied the predicted SNRs differed between the two cases. Anti-symmetric predicted SNRs depended more on the amount of jitter applied, and the symmetric predicted SNRs depended more on the time-window. The next step is to use this information, combined with information on subject performances in other tasks, to define how to most appropriately model the data for the individual subjects. [Research supported by NIH/NIDCD: grant RO1 DC00100.]
Harmonicity and spectral sparsity in speech segregation
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Background
Sound created by a periodic process, as in speech and other natural signals, has harmonic structure—Fourier components at multiples of a fundamental frequency (f0). Harmonicity is closely related to the perception of pitch and is believed to provide an important acoustic grouping cue underlying sound segregation. Here we use a novel speech synthesis method to manipulate the harmonicity of otherwise natural-sounding speech tokens, and report the results of speech segregation experiments with such tokens.

Methods
We utilized elements of the STRAIGHT framework for speech manipulation and synthesis [Kawahara (2006), Acoustical Sci. and Tech.], in which a recorded speech utterance is decomposed into voiced and unvoiced vocal excitation and vocal tract filtering. Unlike the conventional STRAIGHT method, we modeled voiced excitation as a combination of time-varying sinusoids. By individually modifying the frequency of each sinusoid (shifting each component by a fraction of the f0, or jittering each component by a random fraction of the f0), we introduced inharmonic excitation without changing other aspects of the speech signal. We also re-synthesized speech using noise excitation, to simulate the effects of whispering.

To assess the effect of these manipulations on speech segregation, we presented listeners with individual words or pairs of concurrent words spoken by different speakers, and asked them to report as many words as they could.

Results
Performance for isolated words was high for both harmonic and inharmonic speech, and somewhat worse for whispered speech. For pairs of concurrent words, we observed a small performance decrement for inharmonic speech relative to harmonic speech. However, this effect was substantially smaller than the decrement for whispered speech relative to harmonic speech. Surprisingly, the deleterious effect of inharmonicity on speech segregation was significantly larger in musicians than nonmusicians, despite the nonmusical nature of the task.

Conclusion
Harmonic speech excitation aids the comprehension of speech amid concurrent talkers, but the benefit of harmonicity is modest, and seems to be most pronounced in listeners with musical experience. The cost of whispering relative to inharmonic speech indicates that an equally important consequence of voiced speech excitation may be spectral sparsity—the presence of discrete frequency components, as are absent in whispering. By reducing spectral overlap between concurrent signals, sparsity may facilitate segregation via other acoustic grouping cues or top-down lexical matching.
Speech Enhancement Effects Robust to Changes in Spatial Position within a Room
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Background
Previous studies have shown that speech intelligibility in a reverberant room is enhanced by prior listening exposure to that room. Here, this speech enhancement effect is examined under conditions where a carrier phrase and target are presented at different spatial positions in a reverberant room environment that was simulated using virtual auditory space techniques. If the speech enhancement effect relies primarily on the decay pattern of later reverberant energy and not on the specific spatial/temporal patterns of the early reflections, then the effect should be minimally affected by spatial position changes within the room.

Methods
The carrier phrase and target were convolved with separate binaural room impulse responses from different spatial positions in the same room environment. This manipulation altered the temporal positioning of the early reflections but maintained similar patterns of decay in the late reverberation. Intelligibility enhancement data are compared to previous work in which the late reverberation patterns were altered between the carrier phrase and the speech target.

Results
A consistent speech enhancement effect was observed even when spatial position was changed between carrier and target, which suggests that the effect is robust to the resulting changes in the spatial/temporal patterns of early reflections. Consistent with previous work, the magnitude of the effect varies with the amount of room reverberation.

Conclusion
The speech enhancement effect is independent of the specific positioning of the speech source and its carrier phrase in a reverberant room environment.
Remote-frequency masking in school-aged children and adults: Effects of spectral proximity and masker bandwidth
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Background
Previous studies have shown that infants (Werner & Bargones, 1991) and 4- to 6-year-old children (Leibold & Neff, 2011) are susceptible to masking in the presence of a remote-frequency band of noise. In these studies, a 4000-10000 Hz noise band produced significant masking of a 1000-Hz signal in younger listeners, but not in older children or adults. This study examined the influence of envelope modulation rate, masker bandwidth, and spectral separation of the signal and masker on child-adult differences in remote-frequency masking.

Methods
Listeners were 5- to 6-year-olds, 7- to 9-year-olds, and 19- to 30-year-olds (adults). Detection thresholds were measured for a 500-ms, 2000-Hz pure-tone signal in quiet or presented simultaneously with a band of noise, filtered in one of four frequency configurations: 425-500 Hz, 4000-4075 Hz, 8000-8075 Hz, or 4000-10000 Hz. Maskers were played at an overall level of 60 dB SPL, gated on in each interval of a 3IFC adaptive procedure. Two or three estimates were obtained in each condition for each listener. Masking was computed for each listener in each condition, by subtracting the quiet threshold from the masked threshold.

Results
Masking was positive on average for all listener groups in all maskers. However, significantly greater masking was observed for the 4000-4075 Hz masker than for the other three maskers. Remote-frequency masking was comparable for narrowband maskers two octaves above (8000-8075 Hz) and below (425-500 Hz) the signal. There was a modest trend for greater masking in the younger group of children than in the older group of children or in the group of adults, although there were substantial individual differences within listener age groups. Supplemental data were obtained from five additional adults with psychoacoustic listening experience. Following practice, five to seven estimates of threshold were obtained in each condition. In contrast to data obtained from children and naïve adults, results for experienced adult listeners showed little or no masking for any of the four masker conditions.

Conclusion
These results are consistent with those of other studies, showing a reduced ability of children to listen in a frequency-selective manner. These findings also suggest that spectral proximity of the signal and masker plays an important role in masking, even when the signal and masker do not overlap in the auditory periphery.

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Assessing relative sound localization abilities of human listeners in noise
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Background
The 'hemifield code' proposes that the perceived location of a sound source is a consequence of the relative activity of two populations of neurons tuned to contralateral and ipsilateral space respectively. In contrast, the topographic model proposes that neurons exist tuned throughout auditory space. Attempts to disambiguate these two models using brain imaging found that sounds moving away from the midline elicit a greater neural response than those moving toward the midline for broadband and ITD-only stimuli, supporting the hemifield model (Salminen et al 2009, Magezi & Krumbholz, 2010). However, it remains unknown whether these neural response differences result in a behavioural difference in discriminating a change in sound location for outward- versus inward-moving stimuli. We performed a human psychophysics experiment using free-field stimuli to investigate this.

Methods
Subjects made a relative sound localization judgment of stimuli composed of 6 15 ms, white-noise pulses, presented at 10 Hz. Subjects sat within a ring of 18 speakers arranged at 15° intervals spanning 255°. On each trial, the first 3 pulses (reference) were presented from one of 10 locations and the second 3 pulses (target) from an adjacent speaker. The subject made a left/right response to indicate the position of the target relative to the reference. This pulse-train was embedded in noisy background, with independently varying noise being presented from each speaker location. An ITD paradigm with low-pass filtered pulses and an adaptation paradigm with 8-13 pulses at the reference were also tested.

Results
The accuracy of relative sound localization varied throughout auditory space. Relative sound localization abilities were substantially worse in the periphery compared to frontal space, with subjects' best localization performance for judging the relative location of a target sound occurring close to, but not across, the midline. The accuracy of relative sound localization also varied with SNR, with a lower SNR decreasing accuracy. Statistical analysis demonstrated that target location and SNR significantly influenced accuracy, but whether the target sound moved inward or outward did not. The pattern of results observed was very similar in the ITD condition and in the adaptation condition. We compared our experimental data to modeled data based on the models of sound localization outlined above. Relative sound localization abilities were most closely predicted by the two-channel model.

Conclusion
Relative sound localization abilities are affected by reference and target sound locations and signal to noise ratio.

Stimulus Burst Duration Influences Audiovisual Spatial Binding
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Background
Binding, the process by which information from multiple senses is combined to form a unified percept, occurs with a likelihood that is influenced by the spatial discrepancy between auditory and visual stimuli. Previous studies have reported conflicting results indicating that binding likelihood is 50% for audio-visual stimulus discrepancies from as low as 4 degrees, to greater than 15 degrees. One potential source of the disagreement between these studies is differences in burst duration of audio-visual stimuli; studies reporting smaller binding windows used longer stimulus bursts.

Methods
Young adult subjects were asked to localize the source of a continuous train of broadband noise bursts in the presence of a punctate visual distractor that was temporally synchronized, but spatially disparate. Spatial discrepancies ranged from 0° to 20° visual angle, and burst duration was either 50 ms or 200 ms for each trial, with a constant 2 Hz stimulus presentation rate. Auditory localization was measured for each subject and binary logistic regression was used to estimate the audio-visual spatial discrepancy at which binding likelihood was 50%. This metric provides a reliable estimate of the spatial range over which audio-visual binding is likely to occur, since it does not rely on oversampling particular spatial discrepancies or auditory stimulus locations in a manner that could bias subject responses.

Results
Our results demonstrate that increasing audio-visual stimulus burst duration from 50ms to 200ms reduces the audio-visual spatial discrepancy at which binding likelihood was 50%. Additionally, we discovered considerable variability between
subjects, suggesting that there are broad individual differences or unidentified factors contributing to audio-visual spatial binding.

**Conclusion**
Increasing stimulus burst duration reduces the range of audio-visual spatial discrepancies over which stimuli are bound, suggesting that multisensory binding improves with integration over time. These results help explain differences across previous studies.

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**Background**
Studies of sound localization often show substantial differences in performance across participants. Previous studies have suggested that interindividual variability could be related to acoustical differences in localization cues such as the “spectral richness” of the HRTFs (Head Related Transfer Functions). A low spectral richness would limit sound localization performance. In the current study, we investigated the hypothesis that spectral richness can account for interindividual variability in sound-localization performance.

**Methods**
In this study, 25 young (<35 years old) normal-hearing listeners performed an absolute sound-localization task with individual and non-individual HRTFs. The task involved localizing 119 targets distributed uniformly on the surface of a 1.40 m virtual sphere. Participants gave their responses by pointing on a smaller-scale replica of the virtual sphere (God Eye Localization Pointing technique). To reduce procedural-learning effects, participants received procedural training prior to the localization tests. Unsigned left/right and up/down errors, and front/back and up/down reversals rates, were measured as well as the spectral richness of the HRTFs.

**Results**
Large interindividual variability was observed for each of these measured variables. No correlation was observed between the spectral richness and the measured variables. However, the results suggest that interindividual variability in localization performance in the left-right dimension could stem from interindividual variability in the use of the different types of cues available for sound localization in this dimension (binaural/spectral cues). Besides, interindividual variability in sound-localization performance in the up-down and front-back dimensions could be linked primarily to spatial attention and to its variation across the area of space.

**Conclusion**
Interindividual variability in sound localization performance appeared to be more likely related to perceptual origins than acoustical origins.
Effect of Head Movement on Sound Localization Accuracy in the European Starling (Sturnus vulgaris)

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Background
In sound localization experiments, long stimulus durations represent closed-loop conditions that allow the sensory feedback to modify the orientation response of the subject during stimulus presentation. This may yield better thresholds compared to short stimulus durations that represent open-loop conditions and do not allow for orientation responses during stimulus presentation. A previous Go/NoGo experiment on the Minimum Audible Angle (MAA) in the European starling (Sturnus vulgaris) showed that the starlings reach a better MAA for stimulus durations of 1s compared to 100ms in broadband noise conditions (Feinkohl & Klump 2012). For 2-kHz tones, the improvement was not significant. In the present study, we evaluated whether head movements of the starlings during the stimulus presentation in the MAA experiment affect the localization accuracy.

Methods
We recorded videos of the MAA experiment from top view for broadband noise (BBN) and 2-kHz tone (2kT) conditions. For each standard (range 1-5 stimuli) and test signal in a trial, we determined the starling’s head orientation in 9 video frames for the 1s conditions and in 5 video frames for the 100ms conditions, respectively, and calculated the head movement. For occasionally appearing blurry frames that were a result of fast head movement, we only determined the direction of the head movement. Based on these data, we analyzed the effect of the number of head movements on the probability of correct response in relation to stimulus type (100ms BBN, 1s BBN, 100ms 2kT, 1s 2kT). An additional analysis evaluated the role of head movements during the test stimulus presentation only.

Results
Generally, the probability of a correct response increased with the number of head movements. Preliminary data on the analysis of head movements during the test stimuli indicate differences between the stimulus conditions. For stimulus conditions with a duration of 1s, the probability of a correct response increased with the number of head movements, while we found a contrary effect for stimuli with a duration of 100ms.

Conclusion
Head movements allowed the starlings to improve sound localization accuracy in the MAA experiment. While Stimulus durations of 100ms were too short to provide effective head orientation responses, stimulus durations of 1s allowed orientation responses and therefore improved sound localization accuracy.

Rat Cortical Units Display Sharp Hemifield Tuning

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Background
We wish to examine the cortical mechanisms of spatial hearing, specifically of localization and spatial stream segregation. While the cat is a popular model for spatial hearing research, the rodent is a more appropriate model for conducting the in vivo intracellular recordings and pharmacological techniques we envisage. Thus, as a first step, we have developed an anesthetized rat preparation for spatial hearing experiments. Here, we evaluate the spatial tuning of rat primary auditory cortex (A1) to sound-source azimuth with broad-brand noise.

Methods
Extracellular responses were recorded simultaneously from 16 channels of a silicon-substrate probe in field A1 of urethane/xylazine-anesthetized Sprague Dawley rats. Stimuli were 80-ms broad band noise bursts, varying in sound level, and presented from an array of 18 free-field loudspeakers spaced 20 degrees apart throughout 360 deg in the horizontal plane (azimuth). For all cortical units, spatial tuning was characterized by rate-azimuth functions, widths of Equivalent Rectangular Receptive Fields, depths of azimuth-dependent spike-rate modulation, and locations of steepest tuning slopes. Current Source Density (CSD) was used as a functional indicator of the depth of thalamic input layers.

Results
Rat A1 units demonstrated exclusively “hemifield” spatial tuning, with strong responses to contralateral sounds, little or no response to ipsilateral sounds, and a sharp cutoff across the frontal midline leaving little dynamic range to convey information within each hemifield. Moreover, spatial tuning showed surprisingly little variation across sound levels from 20
to 40 dB above threshold. CSD analysis revealed that relative to the putative thalamic input layer, spatial tuning displayed little variation across cortical depth, suggesting that spatial tuning characteristics in rat A1 are already apparent at the level of thalamic input.

**Conclusion**

In contrast to results from anesthetized cats, primates, and ferrets, which display a mixture of contralateral, ipsilateral, and omni-directional tuned neuronal units that expand to increasing suprathreshold sound levels, A1 in anesthetized rats is dominated by neurons that show sharply delimited contralateral tuning with remarkably little variation across increasing suprathreshold sound levels. Previous behavioral investigations have shown that spatial discrimination in rats is excellent across the frontal midline, but that rats cannot discriminate between locations within the lateral fields (Kavanagh & Kelly, 1986). This is in accordance with our results, suggesting that rat A1 units typically signal left or right and do not distinguish among locations away from the frontal midline.

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**Sound Localization in Sagittal Planes: Modeling the Level Dependence**

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**Background**

Sound localization in sagittal planes (SPs), including front-back discrimination, relies on spectral cues resulting from the filtering of incoming sounds by the human body. For short sounds, the vertical range of perceived sound direction depends on the sensation level (SL), an effect often described by the polar gain (linear fit between target and response angles). Data on the polar gain for setups involving both frontal and rear targets seem to be rather limited. Furthermore, models considering the level-dependency are available only as a conceptual description.

**Methods**

Localization responses to brief (3-ms) noise bursts, distributed around normal-hearing listeners both horizontally and vertically in the upper hemisphere, were collected at various SLs from 5 dB to 75 dB in a virtual binaural environment using listener-specific head-related transfer functions. Responses to 300-ms noise bursts presented at an SL of 50 dB were also collected. Localization performance was evaluated in terms of quadrant errors (QE, hemifield confusions), polar angle errors (PE, local root-mean-square errors in the SPs), and polar gain. Performance was also modeled with a localization model based on the comparison of internal sound representation with a template. It consisted of a nonlinear filterbank, a probabilistic decision stage, and a stage retrieving the psychophysical performance parameters. Two template level (TL) conditions were tested: varying with the SL and fixed to 40 dB.

**Results**

For short stimuli, QEs and PEs were lowest at an SL of 40 dB and increased at other SLs. The goodness of fit of the polar gain was overall low, with $R^2$ of 0.058 for SLs $\geq$ 50 dB. The polar response variability was largely level dependent. The psychoacoustic results were accurately predicted by the model with TL fixed to 40 dB and inaccurately predicted by the model with varying TL. For long stimuli, both models were unsuccessful.

**Conclusion**

The level-dependency was confirmed in terms of best performance at an SL of 40 dB. The polar gain did, however, not well describe the effect of SL. Our model was successful in predicting the results for the short stimuli. It failed for longer stimuli, indicating that additional stages such as the MOCR or temporal integration are required. Good results were obtained for the TL fixed to 40 dB, suggesting that the spectro-to-spatial mapping is tuned to a fixed level at the neural level. Funded by the Austrian Science Fund (FWF, P24124-N13).

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**Smooth Pursuit Eye Movements during Tracking of Auditory Targets Suggest Motion-Specific Processing in the Auditory System**

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**Background**

Two conflicting theories address how the perception of auditory motion is perceived: dedicated motion-specific processing, and a strictly position-oriented ‘snapshot theory’. Motion-specific processing posits specialized motion detecting circuits within the CNS, specifically responsive to auditory object velocity. Alternatively, the snapshot theory
assumes sound motion perception is based on periodic sampling of spatial position cues as the target moves through space. Eye movements (saccades and smooth pursuit) may be informative regarding these opposing, though not mutually exclusive, views of auditory motion perception. Smooth pursuit maintains foveal fixation on a moving visual target by smoothly moving the eyes to match target velocity, while saccades ballistically shift foveal fixation from one point to another. Saccades are often employed during visual smooth pursuit when the eyes fall behind and must catch-up to the now peripheral (non-foveal) target. A saccadic strategy of catch-up movements when tracking moving auditory targets argues for the snapshot theory, while smooth pursuit would suggest true motion detection and behavioral tracking. Previous studies have shown little, if any, smooth pursuit of moving auditory targets, but most studies have used high or low velocities that are not ecologically relevant in daily life.

Methods
We presented audio-visual or auditory-only moving targets over a range of realistic velocities (<50º/s) using a concealed loudspeaker/LED target carried on a robotic arm. Eye movements of head-fixed subjects were recorded using electrooculography.

Results
We found that at modest target velocities (20 and 33º/s) smooth pursuit of auditory motion routinely occurs, but with a lower gain (half or less) than for visual targets.

Conclusion
Smooth pursuit of auditory targets occurs, providing strong evidence for a motion-specific auditory processing mechanism. Catch-up saccades are intermingled with pursuit. This suggests that the gain of auditory-driven ocular pursuit is inadequate to accurately track sound objects, requiring a position-dependent correction process co-existing with a true motion processing system.

The acoustical cues to sound location in the adult Guinea pig: measurements of Directional Transfer Functions (DTFs)
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Background
There are three main acoustical cues to sound location, each of which is generated by the spatial-and frequency-dependent filtering of the propagating sound waves with the outer ears: Interaural differences in time (ITD) and level (ILD) as well as monaural spectral shape cues. The guinea pig has been a common model for studying the anatomy, physiology, and psychophysics of binaural and spatial hearing, yet little is known about their acoustical cues.

Methods
Here, we measured the directional transfer functions (DTFs), the directional components of the head-related transfer functions, for 4 adult guinea pigs. DTFs were measured at both ears from 325 locations, with steps of 7.5º in both azimuth and elevation. The resultant localization cues were computed from the DTFs.

Results
The mean weight, head diameter, and the length and width of the pinnae for the guinea pigs were 797 ± 138 g, 30.7 ± 3.5 mm, 32.7 ± 2.9 mm and 10.9 ± 2.5 mm, respectively. In the frontal hemisphere, spectral notches were present for frequencies ~11-19 kHz; in general, the frequency corresponding to the notch increased with increases in source elevation and in azimuth towards the ipsilateral ear. The mean maximum ITD observed across the guinea pigs was 215 ± 5.8 µs. Interaural level differences (ILD) depended strongly on source azimuth and frequency. In general, maximum ILDs were < 12.2 dB for frequencies < 4 kHz, and ranged from 12.2 dB - 40 dB for the frequencies from 4-16 kHz. ILDs for frequencies > ~15 kHz varied with azimuth in complicated ways. Pinna removal eliminated the spectral notches and reduced both ILD and ITD cues.

Conclusion
The acoustical cues to sound location for the guinea pig are consistent with other mammals that have been studied, with respect to their head and pinna size. These results also emphasize the pinna's role in generating these acoustical cues.

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Frequency Dependent Directionality in the Mechanical Response of the Gray Treefrog Ear
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Background
The amphibian ear is inherently directional. Mechanical coupling of the two tympana via the Eustachian tubes and mouth cavity allows each ear to function as a pressure difference receiver. We investigated the frequency and amplitude dependence of this directionality in Cope’s gray treefrog (Hyla chrysoscelis), a species for which there is now a wealth of behavioral data on sound source segregation.

Methods
We measured tympanum vibration with a laser vibrometer in response to short tones (0.60 kHz, 1.25 kHz, 1.625 kHz, 2.50 kHz, and 3.20 kHz) and FM sweeps presented at twelve azimuthal angles (every 30°).

Results
The response of the tympanum to variation in sound source location was frequency dependent. Directionality was most pronounced at frequencies between the two spectral peaks present of the frog's sexual advertisement call (1.25 kHz and 2.5 kHz), with a mean directionality of 6 dB for the 1.625 kHz tone. This frequency dependence was further affected by the volume of air occupying the lungs, with response to lower frequencies most strongly affected by lung air volume. Tympanum response to variation in sound location was sex dependent. Males showed on average 4 dB greater directionality at 1.625 kHz than did females, but tympanum vibration velocities were 3 dB lower across frequencies. Additionally, individuals housed in the laboratory for one year showed similar directionality to frogs recently collected from breeding ponds in the wild, but were less sensitive to high frequency sound (-13 dB at 3.2 kHz).

Conclusion
Our results are congruent with those of previous studies, and represent one of the most thorough investigations to date of the mechanical response of the frog ear. [This work supported by NIDCD R01 5R01DC009582.]

Comparing Two Measures of Absolute Sound Localization Performance in Cats
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Background
Behavioral studies measuring absolute sound localization ability generally implement either an approach-to-target or a pointing task. The approach-to-target task requires the subject to move toward a sound source from a fixed location, while the pointing task typically requires head and/or eye movements toward an auditory target. The fact that subjects usually initiate a head and gaze movement towards the target during the approach-to-target task enables us to measure the accuracy of the initial head and gaze movement and compare it with subsequent speaker selections.

Methods
Cats were trained to localize a broadband noise presented randomly from one of four speaker locations (±30° and ±60°) along the horizontal plane. Trials were self-initiated when the cat's body and head were centered and she was fixating on a central LED. Cats indicated their response by walking to and pressing a lever at the perceived location to obtain a food reward. Reward was delivered only if the first attempt was correct. Trials without a response were classified as “no-go” trials while first selections at the wrong location were "incorrect". A non-categorized measurement of localization accuracy was ascertained by recording the initial head and eye movements immediately following target onset prior to walking to its perceived location. To provide variations in performance, we varied the stimulus duration from 25 to 1000 msec.

Results
Localization performance under both speaker selection and initial gaze movement was very high for long duration sounds but decreased as stimulus duration was reduced. Interestingly, for many of the incorrect and no-go responses, the head and eye oriented to the correct sound source, even when the cat did not approach the correct target. Importantly the eye was significantly more accurate than the head in indicating target location, as the head undershot the eye by as much as 15-20 degrees. At times, the eyes oriented to the targets but the head did not.

Conclusion
Reducing stimulus duration resulted in a systematic decline in both measurements of localization performance. Our data suggest that the head or gaze-orienting response provides a better measure of localization accuracy, especially when the
task is difficult and may minimize bias to specific targets. Apparently, a higher degree of certainty is required to walk to the target, suggesting more involvement in higher-order cognitive processing in this task as compared to gaze orienting.

**Effect of reverberation on acoustic measures relevant for localization (where) and recognition (what) of sounds located at various azimuths and distances: a study of humans and rabbits.**

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**Background**

Binaural room impulse responses (BRIRs) describe how a sound source is transformed into signals in the two ears of a listener located in a reverberant environment. Measures such as reverberation time (T60), direct-to-reverberant (D/R) energy ratio, interaural correlation coefficient (IACC) and acoustic modulation transfer functions (MTFs) can be derived from BRIRs (for MTF, see Schroeder, 1981). Acoustic MTFs have been shown to predict speech intelligibility (Houtgast and Steeneken, 1985). Although BRIRs have been reported in the literature, descriptions of the above acoustic measures are sparse. The goal here is to provide BRIRs of humans and rabbits with a sound source at various locations in two different reverberant environments and to derive the above measures that are relevant for localization and recognition of sounds. This will delineate acoustic conditions that the auditory system faces to accomplish the tasks of localization and recognition in reverberant environments.

**Methods**

We measured BRIRs in an anechoic, moderately- and highly-reverberant environments with a sound source at 0° elevation over a range of azimuths (±180°) and distances (10 cm or 14 cm to 160 cm; Kim et al., 2010; Kuwada et al., 2010). The above acoustic measures were derived from BRIRs for octave-band noise sources with center frequencies of 0.25 ~ 10 kHz.

**Results**

Results for human and rabbit were qualitatively similar. The T60 was a nonmonotonic function of frequency. The values of T60 averaged across frequency were 0.91 and 2.3 sec in the two reverberant environments. With decreasing distance, the D/R increased because the reverberant energy remained approximately constant while direct energy increased. The D/R also depended on azimuth and frequency. The IACC decreased with increasing distance and frequency. Across azimuth, the IACC was large and interaural time difference (ITD) was small when the sound was in the front or back whereas the IACC was small and the ITD large when the sound was lateral. Modulation loss increased monotonically with increasing modulation frequency and distance. When the sound was in the front, modulation loss was large for low carrier frequencies whereas, in the opposite ear, the dependence of modulation loss on the carrier frequency was complex.

**Conclusion**

This study delineates acoustic conditions that the auditory system faces to localize ("where") and recognize ("what") sounds in two different reverberant environments.

**Effect of reverberation on neural coding of sound location (where) and pattern (what) in the inferior colliculus (IC).**

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**Background**

The major functions of the auditory system are to localize and recognize a sound. Traditionally, neural and behavioral studies have examined these two functions separately using artificial conditions that do not represent transformations created by the head and body and environmental reverberations at different sound distances. Furthermore, it is rare to find a study that examined the differences between binaural and monaural stimulation. The goal here is to study simultaneously neural coding of sound location and pattern to binaural and monaural stimulation in environments with different degrees of reverberation.

**Methods**

We measured binaural room impulse responses (BRIRs) in anechoic, moderately- (T60 = 0.91 s) and highly-reverberant (T60 = 2.3 s) environments with a sound source over a range of azimuths (±180°) and distances (10 - 160 cm). Our sound
source was an octave-band noise with center frequencies between 0.25 and 14 kHz that were sinusoidally amplitude modulated (2 - 512 Hz). We created virtual auditory space (VAS) stimuli by convolving these sound sources with the individual animal’s BRIRs. We applied these VAS sounds while recording from neurons in the IC of the unanesthetized rabbit. The center of the noise band was based on the neuron’s best frequency.

Results
In all three environments, we found 4 types of neurons: 1) those that code both azimuth and envelope, 2) those that code only azimuth, 3) those that code only envelope, 4) those that code neither. In all environments, to binaural stimulation, ~25% of the neurons coded only envelope, ~17% coded only azimuth. In anechoic ~55% coded both azimuth and envelope whereas this decreased to ~32% in reverberation. Compared to binaural stimulation, the proportion of neurons that coded envelope to monaural stimulation decreased slightly whereas those that coded azimuth decreased substantially. In general, reverberation tended to degrade azimuth and envelope coding, especially at far distances. This degradation could be almost eliminated by decreasing the sound source distance. In contrast, some neurons were resistant to reverberation even at far distances. Such neurons showed high neural modulation gain.

Conclusion
The IC contains different types of neurons that code azimuth and envelope in combination or separately. The neurons that coded only azimuth or envelope may be the basis for the prominent specialization seen in the non-primary regions of the auditory cortex (Malhotra and Lomber, 2007). The mechanisms that underlie azimuth coding generally require binaural processing whereas the mechanisms that underlie envelope coding generally do not.

Investigating adaptation in the barn owl with a double-stimulus paradigm: a behavioral approach
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Background
During hunting barn owls attend to sounds - as for example rustling generated by prey. The birds typically do not attack upon hearing the first sound, but wait for a second sound. This situation was mimicked with a double-stimulus paradigm.

Methods
It was tested behaviorally whether and how a first or reference sound influenced the head turning of the birds toward a second sound or probe. The inter-stimulus interval between reference and probe was varied between 100 ms and 3200 ms on a log2 scale. Additionally, the owls were stimulated with single stimuli.

Results
Preliminary data collected with three adult barn owls indicate a reduction in precision of head-turning responses when two successive stimuli were presented compared with a situation where only a single sound was presented. Furthermore, head-turning latencies were increased in the double-stimulus condition. This indicated response adaptation. Latencies as well as head-turning precision returned to the level of the single-sound condition if the duration of the inter-stimulus interval was increased. The time constant of recovery from adaptation coarsely matched time constants that were determined in electrophysiological experiments.

Conclusion
Thus, a first stimulus rather lowers than facilitates the response to a second stimulus under the conditions examined.
Efficient Binaural Sound Localization Model for Mobile Agents in Reverberant Environments

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Background
Binaural sound source localization in home environments is encumbered in the presence of strong reverberations. Our goal was to develop an efficient real-time algorithm that improves the localization performance in echoic environments and is able to track sound sources through time and space. In our case, efficiency is required because of the limited processing power of mobile agents.

Methods
As localization cues we used a combination of interaural time differences (ITD) and level differences (ILD), which were analyzed for their reliability using interaural coherence (IC), or the normalized cross-correlation coefficients. A pair of ITD and ILD values is determined for each sample of the digital input signal in each frequency band, and subsumed in a probability density function (PDF) weighted with the corresponding IC values. To merge ITD cues across frequencies we used a summed average in combination with the frequency-specific reliability index. Peaks in the ILD PDFs are matched with our reference ILD values, and the related sound source directions are compared to the peaks in the ITD PDFs to find the most likely sound source locations. The results are integrated over time, to emphasize consistent sound sources and to take gaps in the signal into account.

Results
To evaluate the performance of our algorithm, tests with noise and speech signals in different echoic conditions, and with several degrees of background noise, have been performed. We could show a significant increase of localization accuracy in reverberant environments in comparison to algorithms purely relying on ITD and ILD values.

Conclusion
We combined several simple methods to extract sound source directions with the least possible computational costs and show that this algorithm is still able to extract usable localization information in reverberant, and other adverse acoustical conditions.
Rapid recalibration of auditory distance perception in reverberant environments
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Background
Reverberation and the received sound level provide the most robust cues for auditory distance perception. Specifically, both the direct-to-reverberant energy ratio and the received level increase as the source-to-listener distance decreases. However, the relationship between reverberation, level, and distance varies from room to room. Thus, the auditory system must recalibrate in order to correctly map the cues to distance. Previous experiments showed that listeners are able to "learn" reverberation in a fixed room over the course of multiple hours and days [Kopco et al. (2004) "Learning to Judge Distance of Nearby Sounds in Reverberant and Anechoic Environments." Congress CFA/DAGA; Kopco et al. (2011). "Learning of reverberation cues for auditory distance perception," 161st ASA meeting].

Methods
Current experiments examined the process of rapid adaptation of distance judgments in a small reverberant room during a single 1-hour session. Each session consisted of 8 runs of 80 trials. Presentation level was either fixed during a run, resulting in natural variations in received level with the source distance, or roved by 12 dB for each trial in a run, eliminating the received level as a distance cue. The runs with fixed and roved level alternated during a session. Two groups of subjects participated, differing only by whether they started with a fixed-level or a roved-level run. No feedback was provided. Performance was evaluated in terms of correlation coefficients, means, and variance in responses. Only subjects whose performance was not correlated with presentation level during the roved-level trials were considered in the analysis.

Results
Results depended on whether the level cue was available during the initial run. Correlation coefficients improved over time only if the initial run was performed with roved level. Most of this improvement occurred immediately after the first run. In contrast, subjects who initially had both reverberation and level cues available did not improve over time, and their performance remained inferior compared to the initial-roved-level group throughout the whole session.

Conclusion
These results confirm that rapid recalibration processes occur in spatial perception when listeners enter a new acoustic environment, resulting in improved performance over time. However, the adaptation processes are not triggered automatically, and their occurrence might be conditioned on various factors. For example, distance recalibration occurs only if the level cue is unavailable during the initial exposure to sounds in the new environment.

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Accommodating to New Ears: The Effects of Sensory and Sensory-Motor training
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Background
The auditory system of adult listeners has been shown to re-calibrate to spectral cues altered using in-ear molds (Hofman et al.1998 NatNeuro 1:417-421). Molds initially degraded localization performance but significant improvement followed chronic exposure of 10-60 days (Carlile et al, 2007 ProciCA:221-218). While the functional plasticity of the mature auditory system is itself remarkable, individual differences in the extent and rate of accommodation suggest a number of factors driving accommodation. Prompted by work on cross-modal learning, we hypothesized that it would be possible to facilitate this process by providing multi-modal and sensory-motor feedback during accommodation.

Methods
Using head pointing, 15 subjects localized broadband noise bursts (150 ms) or monosyllabic words from 58 locations roughly equally distributed about an imaginary sphere in a darkened, anechoic environment (5 repeats - baseline). Silicone molds were then fitted, filling approximately 40% of the concha, and localization performance retested (3 repeats). Molds were worn for nearly all waking hours over a period of 10-11 days and performance was tested daily (except weekends) using broadband noise. Subjects were divided into three groups: (1) Control group received no feedback; (2) Visual feedback group: An LED colocated with the sound source, indicated location following each trial; (3) Audio-Visual Sensory-Motor (AVSM) feedback group: In addition to the LED, the sound source pulsed at a rate proportional to the localization error. Subjects were encouraged to explore the stimulus location to minimize localization
error. On the final day of accommodation subjects localized broadband sound and speech stimuli (5 repeats each) with molds and without molds (post-accom).

**Results**

Performance was assessed using the spherical correlation (SC) between actual and perceived stimulus locations and front-back confusion (FBC) rate. Initial mold insertion produced a significant performance reduction. Following accommodation, the FBC rate reduced somewhat for the control group with little change in the SC. Significant improvements were seen for both parameters with both experimental groups, with the largest improvement in SC demonstrated by the AVSM group. Accommodation using the noise stimuli generalized to the speech stimuli.

**Conclusion**

In addition to the accommodation provided by chronic wear, relatively short periods of training involving sensory-motor feedback significantly improved performance that generalized across different stimuli. This has implications for adaptive training to altered auditory inputs.

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**Measuring binaural sensitivity in normal hearing children using headphones**

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**Background**

Motivated by the fact that children with bilateral cochlear implants (CI) show significantly worse sound localization skills than their normal hearing (NH) peers; we investigate whether this can be attributed to poor binaural sensitivity, or other factors. When locating sounds, NH listeners use interaural level differences (ILDs), and interaural time differences (ITDs) in the low-frequency fine structure and high frequency envelopes. The challenge when comparing NH and CI listeners is to use comparably degraded binaural cues. This study is the first to systematically test binaural sensitivity in NH children using CI simulations, and to demonstrate that binaural sensitivity is task-dependent.

**Methods**

Ten NH children (ages 8-10) participated in two experiments. In Experiment 1, intra-cranial lateralization was measured for three stimuli: (1) Spondaic words, (2) Transposed tones (4 kHz carrier modulated at 125 Hz), and (3) vocoder using 500 ms Gaussian-enveloped tone (GET) pulse train (4 kHz center frequency, 100 pulses per second). ITDs (0, ±50, ±100, ±200, ±400, and ±800µs) and ILDs (0, ±1.5, ±3, ±6, ±9, and ±15dB) were imposed on the stimuli and tested in separate blocks of trials. Prior to beginning Experiment 1, children were trained on the task for each set of stimuli. In Experiment 2, just noticeable differences (JNDs) were measured for the transposed tone and GET.

**Results**

When using binaurally degraded stimuli, preliminary results show children have JNDs ranging from ~10-70 µs for ITDs, and ~0.5-2 dB for ILDs. This is comparable to the range of JNDs found in adults. In contrast, on the lateralization task, children perform significantly worse than adults. Although children tend to use the same range of ITD and ILD values to indicated perceived stimulus location, their responses have greater variability across trials, suggesting either task difficulty or spatially ill-defined perception of sound images.

**Conclusion**

The novel finding in this study is that NH children showed binaural discrimination JNDs comparable to adults, suggesting that binaural sensitivity is mature by 8 years of age, even with degraded stimuli. However, the ability to utilize binaural cues for identifying perceived intracranial locations is immature in NH children. Because lateralization is proving to be a more challenging task than discrimination, it may be a better task for assessing binaural maturity. This work offers a benchmark for future studies on binaural sensitivity in children with bilateral CIs.

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Effects of target and masker spatial continuity on speech intelligibility
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Background
In a digit recall task, listeners showed improvement for listening conditions in which a target talker remained stationary and the voice remained unchanged, relative to the case in which the target location and target voice were varied across successive digit presentations (Best et al., 2008). Moreover, recall improved with each successive digit in a sequence. This pattern of performance was attributed a listener’s ability to establish, and refine, a spatial filter over time when continuous auditory stimulation occurred at the filter location. However, it was not clear if changing the spatial locations of the interfering talkers would have a similar impact performance as had been seen with the nonstationary target. The current experiment was designed to address the effect of changing masker and/or target location and pitch on a speech intelligibility task.

Methods
Eight listeners (19-26 years) participated in the experiment. On each trial, listeners seated in quiet rooms heard a target phrase that was created by concatenating 2, 3, 4 or 5 monosyllabic words (Kidd et al., 2008). Two same-sex interfering talkers, also comprising of the same number of monosyllabic words as the target talker, were presented along with the target phrase. The target and two interferers were presented at three of seven possible spatial locations using individualized head-related transfer functions. The spatial location and/or pitch of the target and masker phrases remained fixed or varied word-by-word, resulting in 10 unique listening conditions; all listening conditions remained fixed within a block of trials.

Results
As in Best et al., 2008, word intelligibility improved as a function of number of words contained in the target phrase when the target was stationary and fixed in pitch, and changing either target location or target voice counteracted those improvements. The decline in performance was observed even in conditions when the maskers were stationary and fixed in pitch, suggesting that listeners were unable to utilize masker information to improve performance in the task. And while changing the pitch of the masking talkers did not lead to any further decline in performance, there was a further decline in performance when the maskers changed in spatial locations and varied in pitch.

Conclusion
Data from the current study mostly supports a Listener-Max strategy because performance improved over time when the target was fixed in location or voice. A simple weighted model accounted for the results obtained in the current experiment.

The Effects of Interaural Frequency Mismatch on Spatial Release from Masking
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Background
In normal hearing (NH), binaural benefits occur when frequency-matched inputs from the two ears arrive at the brainstem. Many profoundly deaf individuals receive bilateral cochlear implants (BICls) in an attempt to restore the benefits of binaural hearing, but their performance in noisy environments and ability to localize sound sources is still worse than that of NH individuals. This may be caused by mismatched frequency information between the ears, due to a difference in the insertion depths of the two electrode arrays. Previous studies in which binaural mismatch was deliberately introduced in NH listeners with band-limited acoustical pulse trains, or BiCl users with binaural electrical stimulation, resulted in degraded interaural time and level sensitivity. Without reliable localization cues, it is unlikely that these listeners would be able to make optimal use of the spatial separation from multiple sound sources. The aim of the present study was to investigate the effects of interaural frequency mismatch on speech understanding in a simulated noisy environment.

Methods
Fifteen NH listeners participated in this study. Subjects listened over headphones to five-word sentences spoken by a frontal female target, in the presence of two male maskers speaking IEEE sentences. Maskers were either collocated or 90o to the right of the target. Spatial separation was simulated by filtering stimuli through head-related transfer functions prior to vocoding with an eight-channel sine vocoder. Interaural frequency mismatch was introduced by changing the sine carriers. Listeners responded by identifying the five-word sentence from a closed set. Percent correct scores were
calculated at four signal-to-noise ratios (SNRs) in the two spatial conditions and spatial release from masking (SRM) was calculated by subtracting the percent correct in the separated condition from the corresponding collocated condition.

**Results**

Percent correct scores decreased significantly with increasing mismatch and SRM increased with decreasing SNR. However, there was no interaction between SRM and mismatch.

**Conclusion**

Mismatch reduced speech understanding, but had little effect on SRM. This may be due to access to ILD information at mismatched conditions, which has been shown to be fairly robust to interaural mismatch. It is also possible that limited experience of NH listeners to vocoded speech may have rendered the effects of spatial cues less significant. In contrast, most BiCl users have a prolonged period of adaptation to electrical stimulation, whereby speech understanding improves, and interaural frequency mismatch might still be an important factor for SRM.

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**Word Recognition Benefit from Attention Systems in Older Adults with Hearing Loss**

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**Background**

Speech recognition in noise depends on the integrity of auditory and attention systems to adapt to adverse listening environments. The engagement of the cingulo-opercular error-monitoring system appears to support task performance (Dosenbach et al., 2007; Weissman et al., 2006) and predicts the likelihood of correct word recognition on trials following an error (Vaden et al., 2012). Because the engagement of error-monitoring (Harris et al., 2009) and cross-modal neural systems increases with age (Kuchinsky et al., 2012), we examined the extent to which activity in cingulo-opercular and visual cortices supports correct word recognition for older adults with hearing loss.

**Methods**

In a functional MRI study, 29 older adults (M =67 years; 9 male) with mild to moderately severe sloping sensorineural hearing loss (HL) listened to and repeated 120 consonant-vowel-consonant words presented in multi-talker babble (82 dB SPL) at +3 and +10 dB signal-to-noise ratios (SNRs). Sparse sampling allowed for word presentation and response between image acquisitions. The percent of phonemes correctly reported for each word was tallied. The degree of high and low frequency (HF, LF) HL for each participant was calculated based on weights derived from a normative sample (Eckert et al., 2012) and were independent for these participants.

**Results**

Elevated cingulo-opercular activity associated with incorrect responses predicted better performance on the next trial in the more challenging + 3 dB SNR condition (peak threshold p < .01 uncorrected, cluster extent p < .05 corrected). However, this benefit was reduced with increasing HFHL or LFHL. Whole-brain analyses revealed that visual cortex engagement following an error predicted better subsequent performance only for individuals with greater HFHL.

**Conclusion**

The results demonstrate that older adults engage the cingulo-opercular error-monitoring system, but its efficacy for adapting to error in challenging listening conditions is diminished with increasing hearing loss. For individuals with HFHL, visual cortex activity associated with making an error predicted subsequent correct word recognition, suggesting these individuals employ a cross-modal system to support word recognition. These findings highlight the importance of attention systems that support word recognition and suggest they may be used as biomarkers of hearing loss intervention outcomes.

Work supported by NIH.
Developmental Trends in Listening to Adult and Child Speech in the Presence of another Talker

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Background
An interesting and important question concerns the extent to which children and adults differ in their abilities to hear speech in the background of other sounds, and how children’s’ abilities change as they develop. In this study we investigated the impact of child and adult interfering talkers on children’s ability to focus on child and adult target talkers.

Methods
Target stimuli were of the form ‘Show the dog where the [colour] [number] is’ with the listener required to select the corresponding coloured number from an on-screen response grid. In the conditions with speech maskers, sentences were of the same form as the targets but with a different animal, colour and number. Speech Reception Thresholds (SRTs) were measured adaptively for four conditions: target child speech with masking adult speech, target adult speech with masking child speech, and the same two target speakers in the presence of speech-spectrum-shaped noise matched to the talker. Participants consisted of adults (17 to 51 years) and primary-school-aged children (6 to 11 years).

Results
On the whole, performance for children was very similar to that of adults, except when listening to adult targets in the presence of a child talker. Here, children performed significantly worse than adults, and also showed rapid improvement with age, approaching adult-like levels by age 11. SRTs in the other 3 conditions were close to those of the adults, and showed only small changes across age. The fact that children showed no significant difference in performance from adults for the two target talkers with noise masker suggests it is not the case that children simply find other children’s voices more intelligible than those of adults.

Conclusion
Children, especially those of early school age, find it more difficult to understand an adult talker when the competing sound consists of a single child speaker than when they are listening to a child with an interfering adult talker. In our view, this happens because children are more likely to be distracted by speech from another child than from an adult, and is consistent with the protracted maturation of frontal lobe networks responsible for executive functions related to attention. Such a result has clear implications for listening in the classroom, and unusual distractibility to other children’s voices may be a factor in determining if a child is suspected of auditory processing disorder (APD).
Preparatory and Selective Attention in Children During Multi-Talker Listening

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Background
When differences in the accuracy of multi-talker listening are found between groups of participants, it may be difficult to determine whether the difference results from impaired central attentional control or from a distorted input from the ear. The current project explored a technique for dissociating attentional from peripheral processes that is suitable for use with both adults and children. The technique is based on one established with normal-hearing adults by Hill and Miller (2010, Cereb Cortex 20: 583-590). On each trial, they presented three simultaneous talkers who differed in pitch and spatial location. Importantly, before the acoustic stimuli were presented, participants were cued visually to either the location or the pitch of one of the three talkers. Functional magnetic resonance imaging revealed distinct networks active in preparation for the acoustic stimuli and during the acoustic stimuli. Activity in both networks was influenced by whether participants were cued to the location or pitch of the target voice. Potentially, therefore, brain activity recorded before acoustic stimuli are presented could reveal deficiencies in the control of attention without being confounded by a distorted input from the periphery.

Methods
The current experiment employed a similar paradigm, but recorded the time-course of brain activity using 64-channel electroencephalography. Participants were 22 children 7-12 years of age and 16 young adults 18-27 years of age, all with normal hearing. On each trial, two simultaneous sentences spoken by different adult talkers (one male and one female) were presented from loudspeakers in two spatial locations (one left and one right of fixation). Participants were cued, in advance of the acoustic stimuli, to either the location (left/right) or the gender (male/female) of the target talker. The task was to report key words spoken by that talker.

Results
Event-related potentials (ERPs) differed between trials where participants were cued to the location compared to the gender of the target talker. Differences arose both during preparation for, and during the presentation of, the acoustic stimuli. Children and adults displayed similar ERPs during acoustic stimuli, but showed differences in the preparatory phase.

Conclusion
We conclude that the technique could be used to distinguish preparatory attention from the response to the acoustic input. Ultimately, this paradigm could further our understanding of the contribution of central and peripheral impairments to difficulties with multi-talker listening both in children and in adults.

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Auditory and working memory training for adults with hearing loss: addressing key considerations of outcome measure selection and mechanisms of benefit

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Background
For auditory training (AT) to be an effective intervention for people with hearing loss (PHL), any task-specific learning needs to transfer to functional benefits in real-world listening. This issue has been examined in two auditory training (AT) studies and an ongoing working memory (WM) training study.

Methods
(i) A randomised controlled trial (RCT) of 44 adults with mild sensorineural hearing loss (SNHL) examined the benefits of AT. Participants trained using a phonetic discrimination task in quiet. Measures of speech intelligibility, cognition and self-report of hearing disability were assessed for trained and control participants.

(ii) A repeated measures study identified optimal outcome measures to assess the benefits of AT in 33 hearing aid (HA) users with mild-moderate SNHL. Participants trained on a phonetic discrimination task in noise, thus increasing task difficulty and content validity. Complex measures of cognition (divided and sustained attention, WM and dual-task listening
effort (LE) were assessed pre and post-training. Functional listening benefit was examined using an adaptive two-competing-speaker task (CCRM).

Results
(i) Significant post-training improvements were shown for self-report of hearing, with the largest improvement for a challenging listening situation (p<.05). No improvements were shown for sentences in 8Hz modulated noise (SiMN). Significant improvements were shown for complex measures of cognition (divided attention, p<.01 and WM, p<.05), with no improvements for control participants, nor for simple cognitive tasks.

(ii) Baseline CCRM scores correlated with self-report of hearing (r=.49, p<.01), divided attention (r=.46, p<.05), LE (r=.57, p<.001) and WM (r=.59, p<.01). This supports the contention that competing speech is a more cognitively demanding task than SiMN, which was significantly associated with hearing sensitivity only (r=.39, p<.001). Significant post-training improvements were shown for CCRM (p<.05) and LE (p<.01), with the largest improvements in challenging but achievable task conditions.

Conclusion
Findings suggest outcomes to assess the benefits of AT should be appropriately challenging, yet achievable. Furthermore, benefits of AT may share a greater association with the development of cognitive abilities than the refinement of sensory skills.

A third study directly trains cognition. A double-blind RCT aims to examine benefits of WM training in 54 HA users. Training is performed at an adaptive or a fixed practice-level (active-control). Findings will help inform the most effective training modality (auditory vs. cognitive) for PHL.

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Decoding Auditory Attention (in Real Time) with EEG
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Background
Both magnetoencephalography and electrocorticography recordings have been used to decode which of two competing sources a listener is attending. However, it is not clear whether these techniques might work with Electroencephalography (EEG), particularly in a real-time system. We therefore set out to decode a listener’s attentional focus from EEG signals in real time, knowledge that could be incorporated into next-generation assistive listening devices.

Methods
Offline, we acquired EEG data when a subject listened to a single speech source, from which we estimated a mapping from the EEG data to the perceived speech. The subject then attended to one of two simultaneous speech streams, presented dichotically. The previously estimated system transfer function from the single-source presentation was used to estimate the attended stream in real time. Whichever input speech stream more closely resembled the estimated input was deemed to be the attended stream.

Three decoding methods were tested. The first approach, canonical correlation analysis (CCA), is based on measuring the correlation between audio streams and the EEG signals. The second two approaches estimate the mapping from the EEG to the input stimulus. This can be done using a single channel at a time and summing the result, or by finding a single multi-channel filter that represents all the signals.

Results
We found the best results by estimating a multivariate linear filter that incorporates the channel covariance structure in the least-squares estimation of the impulse response, similar to the approach described by Mesagarani& Chang (2012). Using this approach we could estimate single-speaker data with high accuracy. Notably this approach yielded estimates of the speech envelope that were better correlated with the original speech (r = 0.08) than the other two methods. Applying this to the attention paradigm (after training on single-speaker data), we could predict the focus of attention with 95% accuracy for a one-minute-long sample of dichotic speech. As we shortened the amount of data used to decode, our
accuracy fell almost linearly to about 65% for 10 seconds. Other presentation conditions (i.e., diotic and HRTF) were decoded with lower accuracy than dichotic.

Conclusion
EEG signals can be decoded in real time to determine what natural speech stream a listener is attending with relatively high accuracy.

The Effect of an Enhanced Acoustic Environment on Noise-Induced Tinnitus in Rats
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Background
Noise-induced tinnitus may result from maladaptive plasticity in the brain. Recent studies indicate that enhanced acoustic environments (EAE) can mitigate acoustic trauma-induced hearing loss and tonotopic map reorganization. We aimed to investigate whether an EAE suppresses tinnitus in a rat model.

Methods
Following intense tone exposure (10 kHz, 120 dB SPL, 3 hrs), 12 Sprague-Dawley rats were placed in an EAE consisting of tone pips of random frequency (0.625-20 kHz) and intensity (60-80 dB SPL) for approximately 16 hrs daily over a period of 18 weeks. Another 12 exposed rats were placed in a quiet sound attenuation booth and served as controls. Gap detection (GAP), prepulse inhibition (PPI) acoustic startle reflex behavior, and auditory brainstem response (ABR) were tested on a weekly basis.

Results
Following intense tone exposure, GAP and PPI significantly worsened, indicating behavioral evidence of tinnitus and hearing loss. During the first four weeks, we found gradual improvement in both GAP and PPI data in rats with and without EAE treatments, indicating improved tinnitus. However, continued EAE treatment up to 18 weeks did not improve GAP and PPI results, which was in contrast to the improved GAP and PPI results found in rats without EAE treatment. In addition, the EAE treatment did not induce significant difference in the intense tone-induced ABR threshold shift compared to rats without EAE treatment.

Conclusion
The currently used EAE induced temporary rather than permanent improvement of behavioral evidence of tinnitus in rats. It remains to be investigated whether higher intensity EAE sounds at particular frequency bands will effectively ameliorate acoustic trauma-induced tinnitus.
Noise Induced Hearing Loss on Glutamatergic and GABAergic Gene Expression Changes in Rat Cochlear Nucleus

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Background
Noise-induced damage to the inner ear reduces the neural activity flowing from auditory nerve into the cochlear nucleus. Paradoxically, this leads to a significant increase in spontaneous activity in cochlear nucleus (CN); one of the putative mechanisms for tinnitus. The biological mechanisms that lead to increased spontaneous hyperactivity are largely unknown, but arise from changes in glutamatergic or GABAergic neurotransmission.

Methods
To explore this hypothesis, we unilaterally exposed rats to an intense narrow band noise that caused significant cochlear damage and hearing loss to the exposed ear, but no damage to opposite. Hearing loss was confirmed by auditory brainstem response measurements obtained 14 and 28 days post-exposure. The CN on the exposed and unexposed side of the brainstem were harvested separately at 14 or 28 days post-exposure for qRT-PCR analysis of 84 genes related to glutamatergic and GABAergic neurotransmission.

Results
Unexpectedly, three genes (Gabrg1, Grik4 and Slc17a8) from CN on the unexposed side were significantly up-regulated at 14 days post-exposure. In contrast, two genes (Slc17a6 and Gabrg3) were significantly up-regulated in the CN on the exposed side at 28 days post-exposure. Slc17a6 codes for vesicular glutamate transporter 2 (Vglut2) which is involved in non-auditory glutamatergic synaptic transmission, results consistent with recent work showing increased somatosensory inputs to the CN following cochlear damage. Surprisingly, the CN on the unexposed side showed up-regulation of Slc17a8 which codes for vesicular glutamate transporter 3 (Vglut3), results consistent with recent studies significant changes in Vglut3 in the CN following deafening. Gabrg3, which codes for a subunit of the GABAa receptor gamma 3, also increased in the exposed CN suggesting changes in GABA-mediated inhibition.

Conclusion
Slc17a6 (Vglut2) mRNA expression increased in the ipsilateral CN; this increase may be linked to the invasion of axons from trigeminal nerve. Increased expression of Gabrg3 suggests a compensatory increase of inhibitory, may be occurring to counteract the increases in spontaneous hyperactivity occurring in the CN following cochlear damage.

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Cholinergic modulation of tinnitus related hyperactivity in the Dorsal Cochlear Nucleus.

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Background

Increased spontaneous firing rate (hyperactivity) is observed in various auditory structures in animal models of noise-induced tinnitus. We have previously shown hyperactivity in the DCN (dorsal cochlear nucleus) which undergoes plastic changes and is prominent at frequency loci higher than the frequency of the exposure tone. Previous work has shown that spontaneous firing rate (SFR) in the DCN can be modulated by cholinergic agonists (Chen et al. 1998; Kaltenbach and Zhang 2007). There is also ultra structural evidence of plasticity of the descending cholinergic system after cochlear lesion (Meidinger et al. 2006) which is supported by increased ChAT (choline acetyltransferase) expression in the cochlear nucleus (Jin et al. 2006). In the present study, we sought to examine the functional significance of plasticity of descending cholinergic system in the DCN following noise trauma.

Methods

Adult Syrian golden hamsters were divided in 2 groups, a control group that was not exposed to sound and a sound exposed group (10 kHz, 115 dB SPL for 4 hours). Multi-unit SFR was recorded from the fusiform cell layer (150-250 µm deep from surface) using micropipette electrodes. Effects of the cholinergic agonist carbachol were studied by direct application on the DCN surface. ACSF (artificial cerebrospinal fluid) acted as an internal control and was applied before carbachol application at every recording locus. Long term plots of SFR versus time were obtained from each locus during ACSF and carbachol application respectively. The effect of carbachol on SFR was quantified as mean rate change (mean SFR during carbachol - mean SFR during ACSF / mean SFR during ACSF).

Results

In controls, carbachol had a powerful excitatory effect on SFR at both high (10 and >10 kHz) and low (<10 kHz) frequency loci. This excitatory effect was abolished by atropine. In contrast, carbachol had only a modest effect on thresholds at characteristic frequency in control animals. In exposed animals (short -2-3 weeks and long -6-7 weeks recovery), carbachol had a slight excitatory effect in the low frequency region (short recovery only), but a powerful suppressive effect was observed in the high frequency region. This suppressive effect was blocked by atropine.

Conclusion

Our findings reveal that activation of the cholinergic system can suppress SFR in the high frequency region of the DCN of sound exposed animals. Activation of the granule cell - parallel fiber - cartwheel cell pathway may be a possible route through which the suppressive effect is mediated.
Auditory brain stem responses after noise damage predict tinnitus

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Background
Tinnitus is the phantom perception of a ringing sound in the absence of external auditory stimuli. While startle based gap-detection testing is an objective, behavioral method developed to determine the presence and quality of tinnitus in animal models, objective physiological testing methods are needed to evaluate tinnitus. Following temporary threshold shifts in the auditory brainstem response (ABR) induced by over-exposure to a narrowband noise, guinea pigs show diminished gap-induced reduction of startle response suggesting the development of tinnitus. After recovery of ABR thresholds, there is mounting evidence for loss of auditory nerve drive and compensatory central plasticity, reflected in changes in ABR peak amplitudes and latencies. Here, we extend these studies to provide input-output functions for both left and right ears immediately and several weeks after noise exposure.

Methods
Gap detection and ABRs were recorded in 17 guinea pigs. Nine were unilaterally exposed to a 97 dB, ¼ octave noise band centered at 7 kHz for two hours and eight were sham exposed under Ketamine and Xylazine. Tone-evoked ABRs at frequencies at and outside the exposure band were recorded at baseline, immediately and one week after each noise-exposure procedure, and several weeks after the 2nd noise exposure. Gap detection was assessed biweekly. ABRs were grouped by exposure and evidence for tinnitus.

Results
Five of eight noise-exposed guinea pigs showed impaired gap detection indicative of tinnitus in at least one frequency band. ABR thresholds were elevated near the exposure frequency band immediately after the each noise exposure but recovered to normal within 1 week. In the exposed ear, one week after the noise exposure, P1 amplitudes were reduced but later peak amplitudes were maintained or enhanced. By the final ABR measurement (6-8 weeks after exposure), P1 amplitudes in the exposed ear remained suppressed while P3 and P4 amplitudes were significantly enhanced, more so in animals showing evidence of tinnitus. In the unexposed ear, P3 and P4 were suppressed in animals without evidence of tinnitus but enhanced in animals with evidence of tinnitus.

Conclusion
Reduced P1, but enhanced P3 and P4 ABR amplitudes, correlate with behavioral evidence of tinnitus revealing compensatory changes in central pathways, likely from the VCN, that may play a role in tinnitus. Parallel changes in ABR and gap-detection responses over time suggest the potential of the ABR as an objective physiological measure of tinnitus.

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Acute Tinnitus Perception is Characterized by Altered Oscillatory Activity in Rat Auditory Cortex

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Background
Subjective tinnitus is characterized by the perception of a sound without an acoustic source. Magnetoencephalographic studies comparing people with and without tinnitus have identified specific changes in the spectrum of ongoing activity in auditory cortex. The vast majority of investigations into neurophysiological mechanisms underlying tinnitus-related activity in animals have been carried out in anesthetized preps which abolish perception or in passive animals without confirmation of tinnitus perception and without control for attention. In order to investigate the neural correlates of tinnitus perception in individual attending rats, we developed a novel two-alternative forced choice appetitive behavioral paradigm optimized for the simultaneous recording of neural activity.

Methods
A two-alternative forced choice task was developed to minimize head movement within the acoustic field while permitting several seconds of neural activity to be sampled. Briefly, Sprague-Dawley rats self-initiated a trial by nose-poking in a center hole and waiting between 4 and 8 seconds for a bright light serving as a GO cue. Rats were trained to respond to a right or left feeder depending on the background sound played from an overhead speaker. Correct feeder choices were reinforced with a food pellet: left for steady narrow-band noises (NBN, 1/8th octave bandwidth) of randomized center
frequency, right for amplitude modulated broadband noise (modulated 100% at 5 Hz) or Quiet (speaker off). Microwire electrode arrays were implanted in auditory cortex of trained rats. Neural activity was recorded during the behavior on testing days and filtered offline for local field potentials which were subjected to a multi-tapered spectral analysis. Differences between conditions were evaluated using a Jackknife estimated test statistic.

Results
Following salicylate (200 mg/kg IP), but not saline control, rats changed feeders only on Quiet trials indicating a steady phantom sound was present. During tinnitus, significant changes in oscillatory activity were measured as a decrease in alpha (10 – 14 Hz) and an increase in gamma (40 – 55 Hz) bands.

Conclusion
Changes observed in alpha and gamma bands following salicylate corroborate results from several studies of tinnitus perception in humans. Furthermore, oscillatory activity present during NBN but not Quiet trials indicate that aberrant oscillatory activity during tinnitus perception may not be equivalent to true sound perception and that acute tinnitus perception may be generated cortically. We hope to extend this approach to better characterize the neural mechanisms of phantom sound perception in the brain.

Biofeedback as a potential prophylactic treatment for blast-induced tinnitus.
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Background
The American Tinnitus Association has estimated that over 50 million Americans experience tinnitus. Currently the only known effective treatment is the use of biofeedback techniques. Because tinnitus is thought to be perpetuated via a hyper-excitation of the amygdalo-auditory cortex circuits, biofeedback therapy may act to increase GABAergic inhibition of these circuits.

Methods
To test our hypothesis, animals were tested for tinnitus using acoustic startle responses (i.e., startle, pre-pulse, and gap detection tests). They were then exposed to blast sounds (90 dB SPL; 1 hour/day). During this sound exposure, animals became accustomed to the sounds and started to sleep during the exposures. After 2 weeks of sound exposure, animals were exposed to a 20 psi blast pressure wave, and then subsequently tested for tinnitus twice per week post-blast-exposure. The incidence of tinnitus was then compared in the sound-exposed group of animals versus blasted non-sound-exposed animals.

Results
The incidence of tinnitus with sound-exposure (i.e., biofeedback treatment) was substantially decreased when compared to similar animals that did not receive the biofeedback treatment.

Conclusion
We hypothesize that biofeedback therapy may be a simple and cost-effective prophylactic treatment for individuals that are at risk for blast-exposure.

Functional Changes in Amygdala and Striatum in Rats with Noise-Induced Hearing Loss
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Background
Most studies of noise-induced hearing loss (NIHL) have focused on functional and structural changes within the classical auditory pathway. However, other regions outside the classical auditory pathway may also respond to acoustic stimulation, for example the striatum which is involved in vocal motor control and the amygdala which play a role in auditory fear conditioning and can modulate the acoustic startle reflex. To determine how acoustic overstimulation
affected these non-classical auditory regions, we measured neural firing patterns in the lateral amygdala (LA) and the overlying striatum area of rat after inducing a high-frequency NIHL.

**Methods**

Rats were exposed to a narrowband noise (16-20 kHz) at 102 dB SPL. Afterwards, auditory brainstem response (ABR) was measured to determine the degree of NIHL. Multiunit recordings were obtained from the LA and striatum using 16 channel electrodes. Then, the cochleograms were prepared to characterize the location and degree of IHC and OHC loss.

**Results**

The noise exposure resulted in ABR threshold shifts of 30-40 dB at the high frequencies. This was associated with varying amounts of IHC and OHC loss in the basal turn of the cochlea. Neural thresholds in the LA and striatum of the noise-exposed rats were normal at low frequencies, but elevated at the high frequencies, consistent with the ABR data and cochlear lesions. In 2 of the 4 noise-exposed rats, neural responses to suprathreshold stimuli decreased with increasing frequency, i.e., high frequency hypoactivity. However, in the other 2 rats, suprathreshold neuronal responses in the LA, but not the striatum, were hyperactive (enhanced) at the low-frequencies bordering the hearing loss region and hypoactive at the high frequencies within the NIHL. The 2 rats with edge-frequency hyperactivity had fewer missing OHC than those with strictly high-frequency hypoactivity.

**Conclusion**

The frequency-dependent loss of auditory sensitivity in the amygdala and striatum following noise exposure was consistent with the ABR threshold shifts and cochlear lesion. In some noise exposed rats, neural responses in the LA were hyperactive at the transition from normal hearing to impaired hearing region; the hyperactivity at the transition region appears to be related to the survival of OHCs at the edge of the cochlear lesion. In a broader context, the hypoactivity seen in the striatum would likely impair vocal motor control whereas the hyperactivity seen in the amygdala might be related to hyperacusis or emotional reaction to sound.

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**Corticosterone Increases Significantly during Salicylate Induced Tinnitus**

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**Background**

Previous reports suggest that the onset or increase in tinnitus in patients with hearing loss is often associated with periods of high stress. Since high levels of stress could conceivably play a critical role in the development of tinnitus, we hypothesized that the onset of tinnitus might be linked to an increase in the circulating stress hormones. To test this hypothesis, we treated rats with a high dose of sodium salicylate, an extremely reliable method for inducing tinnitus.

**Methods**

Male Sprague-Dawley rats were treated with saline or 50, 150 or 250 mg/kg of sodium salicylate. Blood was sampled at 48 h before and 2, 24 and 48 h post-salicylate. Blood serum corticosterone levels were measured using a competitive ELISA assay kit. Corticosterone measurements were obtained from both anesthetized and un-anesthetized animals treated with salicylate.

**Results**

The 250 mg/kg dose of salicylate produced a massive increase in corticosterone levels at 2 h post-salicylate (614 ng/ml serum) compared to saline controls (25 ng/ml serum). Corticosterone levels decreased as the salicylate dose was lowered. The 50 mg/kg dose had no effect on corticosterone levels whereas the 150 mg/kg dose produced a modest increase in corticosterone. Corticosterone levels were highest 2 h post-treatment and then gradually declined over the next 24-48 h. Corticosterone levels increased in both anesthetized and un-anesthetized animals; however the increases were substantially less in anesthetized animals.

**Conclusion**

High doses of salicylate increased corticosterone levels in a dose and time dependent manner. The 50 mg/kg salicylate dose, which does not induce tinnitus, did not increase corticosterone levels whereas the 150 mg/kg salicylate dose, the lowest dose that induces tinnitus, produced a modest increase in corticosterone. The decrease in corticosterone levels following the 250 mg/kg salicylate dose decreased over time but was still elevated in some rats at 48 h post-treatment. Thus, the recovery of corticosterone levels over time parallels the behavioral recovery of tinnitus. These results are
consistent with the hypothesis that the stress response induced by high doses of salicylate contributes to the induction of tinnitus.

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Histological Analysis of Vestibular Organs in Caspase 3 Deficient Mice
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Background
Caspase 3 deficient (Casp3 KO) mice exhibit circling behavior suggestive of vestibular dysfunction. In previous studies we have shown that Casp3 KO mice have absent horizontal vestibulo-ocular responses (angular acceleration), but have normal otolith-ocular responses (linear acceleration). Our goal was to identify histological characteristics which correlate with the behavioral functional analysis results, in order to determine the mechanism of the vestibular dysfunction in these mice.

Methods
Whole mount vestibular organs of Casp3 KO, Casp3 heterozygous and WT mice, were labeled with hair cell markers (phalloidin, calretinin) and observed at different ages. Younger (0-6 months) and older (7-24 months) age groups were compared. The following metrics were measured in the anterior-, lateral-, posterior ampullae, utricle and saccule: hair cell number, sensory epithelia area, hair cell density, striolar/extrastriolar hair cell density. Two-tailed t-tests and analyses of variance were used for statistical analysis.

Results
In younger ages, the vestibular organs were smaller in Casp3 KO mice, especially in the anterior- and lateral ampullae and utricle (p<0.05), while those of heterozygous and WT mice were normal in size and shape. In Casp3 KO mice, hypomorphisms in the vestibular organs varied in degree, ranging from unilateral, bilateral or normal. Occasionally the anterior-, and lateral ampullae were found to be fused together, or the utricle in narrower shape.
In older ages, we observed fewer hair cells in all vestibular organs and in all genotypes (p<0.05). The most striking observation was that in older Casp3 KO mice (>6 months), the saccular hair cells were almost completely degenerated, while heterozygotes and WT type mice did not have hair cell loss even in older ages. On the other hand, there was no significant hair cell loss in the utricle of older Casp3 KO mice (p>0.05).
Younger heterozygotes had higher hair cell count and density in all organs, but both decreased significantly with age. Heterozygote utricles and saccules both increased in size with age.
There was a trend that hair cell loss occurred mainly in the striolar area rather than the extrastriolar area in all genotypes.

Conclusion
In Casp3 KO mice, the hypomorphic shape and fewer hair cell numbers of the lateral ampulla correlate well with the hypofunctional horizontal VOR. The complete loss of saccular hair cells in Casp3 KO mice may be used as a unique model to differentiate individual otolith organ function.
Distributions of the cVEMPs along the sternocleidomastoid muscles during neck rotation and flexion in normal human subjects

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Background

Tone burst-evoked myogenic potentials recorded from the sternocleidomastoid (SCM) muscle (cervical VEMP or cVEMP) are commonly used clinically to assess the vestibular function. The SCM is not only one of the largest muscles in the neck, running from the sternum and clavicle to the mastoid and occipital bone, but also has multiple functions. When the pair of SCMs contract together, the neck is flexed and the chin is led downward. When a single SCM acts alone, the head is rotated to the opposite side. The cVEMP is commonly recorded at the upper third of the SCM during head rotation or neck flexion. In a previous study, we examined the distributions of cVEMPs along the SCM during head rotation. In the present study, we further compared the cVEMP distributions along the SCM during neck rotation and flexion.

Methods

Surface electrodes overlying the SCM muscles were used to record cVEMPs in 12 healthy adult subjects evoked by tone bursts delivered to the ipsilateral or the contralateral ear (tone burst, 8ms plateau, 1ms rise/fall, 130 dB SPL, 50-4000Hz). Tonic contractions of the SCMs were achieved by turning the head to the right or flexing the head downward.

Results

Preliminary analysis revealed that cVEMPs exhibited differences in amplitude, latency and morphology during neck rotation and flexion. The differences were tone frequency-dependent.

Conclusion

Neck rotation and flexion recruit different innervations patterns along the SCM, which may influence the cVEMPs recorded at different sites along the SCM. Future studies are to characterize these responses and explore their clinical implications.
The Neural Basis of Clinical Vestibular Responses to Bone Conducted Vibration (BCV) and Air- Conducted Sound (ACS)

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Background
New clinical tests of human otolithic function use 500Hz BCV and ACS stimuli and measure vestibular evoked myogenic potentials from surface EMG electrodes beneath the eyes (oVEMPs) or over neck muscles (cVEMPs). A different clinical indicator uses intense 50Hz or 100Hz vibration of the mastoid to induce vibration induced nystagmus (VIN) in patients with unilateral vestibular loss. This study asked are some utricular and saccular afferents activated by both 500Hz BCV and ACS? do these stimuli activate semicircular canal neurons? Is 100Hz BCV also specific for otolithic afferents or are semicircular canal afferents also activated by intense low frequency vibration?

Methods
Single primary vestibular neurons were recorded extracellularly in guinea pigs anesthetized with Ketamine and Xylazine or rats anesthetized with pentobarbital. The neurons were identified by their location and their response to angular and linear accelerations. BCV stimulation was delivered either by a Radioear B-71 bone oscillator cemented to the skull or a Bruel and Kjaer 4810 Minishaker held against the stereotaxic frame. ACS stimulation of up to 140dB SPL was delivered by a TDH-49 headphone. Juxtacellular neurobiotin injections were used to identify whether the receptors synapsing on the recorded afferent, originated from the otoliths.

Results
Many otolithic irregular neurons were activated at low threshold by 500Hz BCV. Many of these were also activated by high intensity 500Hz ACS. Neurobiotin labelling showed that in both guinea pigs and rats these afferents originated from the utricular macula and also from the saccular macula, usually from presumed Type I receptors at the region of the striola. Semicircular canal neurons were not activated by 500Hz BCV or ACS up to the maximum intensity we delivered. At 100Hz otolithic afferents were strongly activated and synchronized with the stimulus, but with 100Hz high intensity vibration stimulation, irregular semicircular canal neurons and even some regular canal afferents were activated and synchronized to the stimulus.

Conclusion
The wide presumption in clinical vestibular testing - that ACS only activates saccular afferents in the inferior vestibular nerve - is not correct: these results show that 500Hz BCV and ACS selectively activate otolithic but not canal neurons. So 500Hz BCV and ACS are effective specific otolithic stimuli for testing oVEMPs and cVEMPs. However at low frequency this specificity is lost; high intensity 100Hz BCV activates both otolithic and semicircular canal neurons. The activation of these canal neurons is likely mainly responsible for vibration induced nystagmus in human patients.

Updating Alexander’s Law and the Oculomotor Neural Integrator

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Background
Alexander’s law (AL) states that the nystagmus slow phase velocity in patients with peripheral vestibular lesions is lower when looking in the slow compared to the fast phase direction. It has been hypothesized to arise either due to adaptive changes in the velocity-to-position neural integrator (NI), or as a consequence of processing of the vestibulo-ocular reflex. It has also been reported to develop with a latency of 20-30 s after start of nystagmus, and proposed that only an abnormal stimulus evokes AL.

Methods
We tested AL in healthy volunteers during unilateral warm and cold caloric stimulation, as well as simultaneous bilateral bithermal stimulation. The latter condition simulates the normal push-pull stimulation of natural head movements. We also tested subjects during long duration, constant angular acceleration, which produced constant nystagmus for > 35 seconds, in order to stimulate the vestibular system with real head rotations.

Results
All caloric stimuli evoked AL, although in < 10% it was inverted. On average, the development of AL did not lag nystagmus. During constant acceleration, i.e. an almost natural stimulus, all volunteers developed a strong and stable AL, corresponding to integrator time constants of < 3 seconds which are much less than normal time constants (~25 seconds).
and similar to those observed in patients with acute unilateral vestibular lesions showing AL. Alexander's law also developed, on average, in less than 10 seconds.

**Conclusion**

We conclude from these experiments, that AL is not an adaptive reaction and that it occurs immediately with natural vestibular stimulation. We propose a model of processing of vestibular and eye position signals in the vestibular nuclei, that explains the immediate start of AL.

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Functional Dissection of the Neural Circuitry Responsible for the Vestibuloocular Reflex in Larval Zebrafish

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**Background**

We have combined a systems-level behavioral characterization with molecular neuroscience to identify and characterize the neural circuitry underlying the vestibuloocular reflex (VOR) in the larval zebrafish. Young zebrafish develop excellent compensatory ocular rotations following body tilts over the first week of life, after which the larval zebrafish can reach mammalian levels of performance. This behavior is mediated by the utricle only, as the semicircular canals are not yet functional. We have shown that a classic three neuron circuit (primary vestibular afferents, central processing neurons, and ocular motoneurons) is necessary for gaze stabilization following body rotations.

**Methods**

We developed a novel transgenic line which expresses in vestibular afferent neurons and second-order central neurons. We used this line to perform laser ablations, optogenetic excitation, single-cell fills, and electrophysiology to characterize the role these neurons play during the VOR.

**Results**

By targeted photoablation, we found that the neurons in this line are necessary for the VOR. Using optogenetics, we can elicit stereotyped eye movements comparable to those observed during body rotation, suggesting sufficiency. To understand how these neurons function as a population, we sought to identify subgroups of neurons likely to subserve specific aspects of the VOR. We build a model of the larval zebrafish VOR circuit, using previous insights from mammalian preparations, and known projection patterns in teleost fish. Our model predicts anatomically distinct classes of second-order central vestibular neurons. We labelled individual neurons in our transgenic line, traced their morphology, and showed that the predicted classes of neurons are indeed present. We are currently using ablation and activation to test the functional role of these specific subpopulations. Finally, we report on the basic electrical properties of central vestibular neurons, identifying both mature and immature classes.

**Conclusion**

Our transgenic line of fish offers molecular-level control over the population of neurons responsible for a dynamic sensorimotor transformation. The anatomical and electrophysiological properties of labelled neurons are compatible with a simple model using stereotyped wiring to link the sensation of gravity to muscle activation to stabilize gaze.
Magnetic field-induced eye movements in mice
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Background
Roberts et al. demonstrated that static magnetic fields induce nystagmus in humans by peripheral vestibular stimulation and that this effect may be mediated by a magnetohydrodynamic force (1). This hypothesis can be tested by changing head orientation within static magnetic fields, but MRI magnet bore dimensions limit the ability to do this with humans. A mouse model would allow greater flexibility in assessing multiple head orientations. The aim of this study was to demonstrate magnetic field-induced nystagmus in mice.

Methods
Wild-type C57BL/6J mice were exposed to a 4.7T Earth-horizontal magnetic field. Prior to exposure, the mouse’s head was secured to a non-ferromagnetic post and positioned so the horizontal semicircular canals were earth-horizontal. Monocular eye movements of the mouse’s right eye were recorded via video-oculography in darkness with infrared illumination. Mice were positioned in the center of the magnet bore for at least 1 minute in each of several head orientations. Slow-phase nystagmus velocity (SPV) was calculated and an exponential decay was fit to the peak SPV over time.

Results
When placed in the bore nose-first, all mice demonstrated robust left-beating nystagmus lasting approximately 20 seconds on first entry. Mean peak SPV of six mice was 282°/s and the time constant was 3.62 s (SD 0.39) over the initial interval within the bore. Beats of nystagmus direction reversal were seen on a few trials after exiting the bore. When placed in the bore tail-first, nystagmus direction reversed from the nose-first orientation. When placed in the bore left-ear-first or right-ear-first, few eye movements were observed.

Conclusion
Mice exposed to high-strength magnetic fields demonstrate robust magnetic field-induced nystagmus that depends on magnetic field orientation. These findings support the hypothesis that static magnetic fields induce peripheral vestibular stimulation through a Lorentz force created by resting ionic currents within the labyrinth. The peak SPV and decay are both faster than in human subjects and may reflect poor velocity storage and relatively weak velocity-to-position integrator in the mouse.

Glycine receptor deficiency and its effect on the vestibulo-ocular reflex: a study on the SPD1J mouse.

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Background
Our recent study in three strains of mice with naturally occurring glycine receptor (GlyR) mutations identified a difference in MVN neuronal action potential (AP) and discharge properties. When compared to wildtype mice, this difference in spastic, spasmodic, and oscillator mice manifested itself as a reduction in background discharge rates, larger AP after-hyperpolarizations (AHP), and lower sensitivity to current steps. We hypothesized that these differences, particularly the reduced excitability of MVN neurons in mutant strains, would impair the slow-phase VOR response. In addition, because glycine is also used by the omnipause neurons (OPN) of the saccadic and quick-phase system we predicted that the quick-phase amplitudes and peak velocities would be altered.

Methods
To study the effects of GlyR deficiency we tested the three dimensional angular vestibulo-ocular reflex (3D VOR) of homozygous (affected) and heterozygous (control) spasmodic mice (13-15 weeks). Slow-phase and quick-phase velocity of the VOR response was analysed for sinusoidal head rotations between 0.1 to 12 Hz reaching peak-velocities of 20, 50 or 100 º/s and for transient acceleration steps of 3000 or 6000 º/s² reaching a velocity plateau of 150 º/s (for 3000 º/s²), 300 and 600 º/s (for 6000 º/s²).

Results
The slow-phase VOR of affected mice was significantly lower during the constant velocity plateau of the transient acceleration step stimuli and at low-frequency (0.1 – 0.5 Hz) high-velocity (100 º/s) sinusoidal stimuli. Compared to control mice, quick-phases in affected mice were reduced in number (30 – 70 % less per cycle, depending on stimulus) and had considerably smaller peak-velocities (38% ± 5% less) and amplitudes (55 ± 5% less).

Conclusion
Glycinergic transmission deficiency in homozygous spasmodic mice predominantly affects the quick-phase system of the VOR. Our results support the notion that glycinergic OPNs in the saccadic/quick-phase system act as a regulator of quick-phase gain rather than as a gate for quick-phases.
Partial restoration of the VOR in humans by motion-modulated electrical stimulation of the vestibular system

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Background
Patients with a bilateral vestibular loss (BVL) suffer from imbalance and oscillopsia. Currently there is no treatment. We explored whether the vestibulo-ocular reflex (VOR) can be artificially restored in BVL patients using motion-modulated electrical stimulation of the vestibular system.

Methods
A unilaterally deaf BVL subject (male, 50 years old) was fitted with a modified cochlear implant (Med-EL®; Innsbruck, Austria) providing an extracochlear “vestibular” electrode positioned in the vicinity of the posterior ampullary nerve. After adaptation to steady-state suprathreshold electrical stimulation, motion sensors were fixed on the subject’s head and used to modulate stimulation currents delivered to the “vestibular” electrode. Eye movements were recorded with a video-oculography system incorporating 6 DOF head motion sensors (EyeSeeCam VOG; Munich, Germany) during whole body rotations about an earth-vertical axis (sinusoidal, 1Hz, 30°/s peak velocity).

The VOR response was tested in two conditions: a) without any electrical stimulation (SYSTEM OFF), and b) with motion-modulated electrical stimulation of the vestibular pathways (SYSTEM ON). Electrical stimuli consisted in charge-balanced biphasic current pulse trains (200µs/phase at 400 pulses-per-second), amplitude modulated via the yaw signal captured by the head-fixed motion sensor (gains: 1.7µA°/s–3.4µA°/s).

Recordings were analyzed on a cycle-by-cycle basis; mean (±SD) responses were calculated over all cycles. VOR gains were computed using sinusoidal fits to raw data.

Results
Control subjects with normal vestibular function presented VOR gains ranging 0.7-1.0 (Fig. 1a). For the BVL subject in the SYSTEM OFF condition, VOR gain was consistently ≤0.1 and eye movements were not correlated to yaw motion data (Pearson’s correlation: r=0.07, p<0.05; Fig. 1b). In the SYSTEM ON condition, the VOR gain of the BVL subject increased progressively as stimulation strength increased and eye movements were highly correlated to yaw motion data (Pearson’s correlation: r=-0.69, p<0.0001). The maximum gain observed was 0.36(±0.18), a substantial recovery towards normal function (Fig.1c).

Conclusion
This is the first demonstration that the VOR can be partially restored in a human BVL patient via motion-modulated electrical stimulation of a vestibular nerve branch. This is a fundamental first step towards the demonstration that such devices could help rehabilitate patients with BVL. Our next step will be to extend these results to other patients implanted with similar devices.
Bilateral Stimulation of the Vestibular Labyrinth in Rhesus Monkeys Using a Multichannel Vestibular Prosthesis
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Background
Maximizing efficacy of vestibular nerve branch stimulation is a key goal in development of a multichannel vestibular prosthesis (MVP) intended to restore sensation of 3-dimensional head movement in individuals disabled by chronic bilateral loss of labyrinthine hair cell function. Unilateral prosthetic stimulation in bilaterally vestibular-deficient (BVD) rhesus monkeys elicits vestibular-ocular reflex (VOR) responses with marked asymmetry in gain during rapid head rotations. VOR gain is lower for head rotations toward the non-implanted side (away from the prosthesis side) than for rotations in the opposite direction. Reducing this gain asymmetry by enhancing contralateral responses would be an important step toward improving gaze stability in BVD individuals using a MVP. Noting that an individual with two normal labyrinths has a much more symmetric VOR than an individual with a single normal labyrinth (e.g., after unilateral labyrinthectomy), we hypothesized that bilateral, complementary prosthetic stimulation of the vestibular labyrinths will elicit responses with less asymmetry than observed with unilateral stimulation.

Methods
We implanted MVP electrodes in semicircular canals of both ears in one rhesus monkey previously rendered bilaterally vestibular-deficient via intratympanic gentamicin administration. We characterized horizontal VOR responses using scleral coils in darkness during unilateral and bilateral prosthetic electrical stimulation.

Results
Bilateral complementary stimulation produced significantly higher VOR gain and improved symmetry compared with unilateral stimulation.

Conclusion
Consistent with a prior study characterizing horizontal VOR performance during bilateral stimulation of the lateral semicircular canals in squirrel monkeys, our results support the hypothesis that bilateral vestibular prosthetic stimulation can achieve better 3D VOR performance in rhesus monkeys than possible with unilateral stimulation. However, this improvement comes at the expense of twice the surgical risk and, barring implant designs with a single processor driving electrode arrays in both ears, approximately twice the cost per patient once this technology becomes a part of routine clinical practice. The relative risks, costs and benefits between bilateral and unilateral stimulation will ultimately depend on the extent to which changes in stimulation timing and waveforms can achieve good symmetry with unilateral implantation.
System Integration of an Application Specific Integrated Circuit to Advance Toward a Fully Implantable Multichannel Vestibular Prosthesis

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Background
The Johns Hopkins Multichannel Vestibular Prosthesis (MVP) has demonstrated success in modulating vestibular afferent fiber firing rates to encode rotational head movements for those with profound bilateral vestibular hypofunction. Current implementation of the MVP follows a semi-implantable approach, with implanted electrodes coupled to an external component including motion sensors, processing circuitry, and the power source. Decoupling of the external component would cause loss of power for the prosthesis and sudden onset of vertigo. To avoid the issue of decoupling, development of the next generation MVP (MVP3) aims for a fully implantable system, where device size becomes a key consideration.

The development of a complementary metal-oxide semiconductors (CMOS) application specific integrated circuit (ASIC) provides the ability to greatly reduce size of the device. The goal of this experiment was to determine whether integration of the CMOS ASIC would offer the same clinical benefits as the existing MVP architecture (MVP2).

Methods
We developed a CMOS ASIC to replace most of the circuitry for the output of the prosthesis, leaving three main components that make up the MVP3: motion sensors, a microcontroller and the ASIC. Integration of the ASIC into the existing MVP architecture required modification of existing software for the microcontroller to supply correct commands to the ASIC based on the motion sensor input to the system. The ASIC was designed to efficiently organize commanded signals from the microcontroller with regard to current amplitude, timing, and polarity.

After bench testing of ASIC integration, the MVP3 system was verified in a rhesus monkey rendered bilaterally vestibular-deficient via intratympanic gentamicin. We rotated the monkey sinusoidally in darkness at 0.1-5 Hz, at 50-200 °/s peak. The 3-dimensional vestibular ocular reflex (3D VOR) responses were recorded and compared to eye movements evoked by the MVP2.

Results
Bench testing of the ASIC with custom microprocessor software allowed for correct encoding of motion to frequency modulation of pulses. Physiological tests produced 3D VOR eye velocity movements that are statistically similar to the MVP2.

Conclusion
Comparison of the MVP3 with the MVP2 showed no significant difference, thereby verifying physiological implementation of the ASIC interface between the microcontroller and electrodes. With these initial tests of feasibility for the ASIC, future development will further integrate circuitry for the microcontroller into the ASIC. This continued miniaturization and reduction of power consumption will provide a solution for a fully implantable prosthesis.

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Heterogeneity in the Population of Vestibular Neurons that are Activated by Sinusoidal Galvanic Vestibular Stimulation (sGVS) and Send Projections to Pre-sympathetic Neurons in the Ventrolateral Medulla

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Background
Vestibulo-sympathetic pathways mediate rapid changes in blood pressure in response to movement and adjustments in head and body position. We previously demonstrated that there are direct projections from the caudal vestibular nuclei to the rostral and caudal ventrolateral medullary cell groups (RVLM and CVLM, respectively), the key pre-sympathetic sites for integration of cardiovascular signals. We have also shown that neurons in this pathway are activated by sGVS. The goal of the present study was to identify the topography and chemical anatomy of these projections.

Methods
Unilateral iontophoretic injections of the retrograde tracer FluoroGold (FG) were placed in the RVLM or CVLM of rats. After 7-10 days, the animals were anesthetized and received sGVS while changes in blood pressure were recorded using photoplethysmography. Animals were perfused 90 min after the stimulation. The brains were harvested and the
brainstems were sectioned serially. Sets of sections were used to map the tracer injection sites, and other series’ of sections were used for multiple label immunofluorescence studies of the caudal vestibular nuclei. These latter experiments used c-Fos protein accumulation as a marker for neurons activated by sGVS, as well as immunolabels to identify the tracer, glutamate, GABA, and imidazoleacetic acid-ribotide (IAA-RP), a neuromodulator involved in blood pressure regulation, in vestibular neuronal somata.

**Results**
FG was present bilaterally in cell bodies of the inferior and caudal medial vestibular nuclei (IVN and MVNc, respectively) that had c-Fos protein accumulation in response to sGVS. The sGVS-activated vestibular neurons with projections to RVLM or CVLM were distributed approximately equally in MVNc and IVN (101 vs 90 cells observed with co-localization of FG and cFos in MVNc and IVN, respectively). Within MVN and IVN, approximately equal numbers of cells projected to CVLM and to RVLM. However, almost twice as many cells in each vestibular region sent ipsilateral as opposed to contralateral projections. These studies also revealed differential localization of glutamate, GABA and IAA-RP immunofluorescence in the MVNc and IVN projections to ipsilateral and contralateral RVLM and CVLM.

**Conclusion**
We conclude that there are multiple topographical and chemoanatomical components of the vestibulo-sympathetic pathway.

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Double Oscillations in Blood Pressure and Heart Rate from Monaural Sinusoidal Galvanic Vestibular Stimulation (sGVS) in the Rat

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Background
Binaural sinusoidal galvanic vestibular stimulation (sGVS) induces oscillatory changes in blood pressure (BP) and heart rate (HR) in anesthetized rats that are associated with vasovagal responses (VVR). sGVS also induces similar oscillations in muscle sympathetic nerve activity (MSNA) in alert humans. There are generally two oscillations of BP and HR in rats and two cycles of MSNA in humans for each cycle of low frequency sGVS. We questioned whether these double oscillations were related to known vestibular mechanisms and could shed light on the processing of sGVS signals from the vestibular nerve to the autonomic system.

Methods
BP and HR were measured with intra-aortic sensors in three rats anesthetized with 2% isoflurane. The labyrinths were stimulated at frequencies of 0.025-0.5 Hz with currents of 3 mA through Ag/AgCl needle electrodes inserted over the external auditory meati on both sides (binaural stimulation) or in front and behind the external auditory meatus of one ear (monaural stimulation). Single sinusoids were also given monaurally and binaurally at 2 minute intervals. In addition, rats were also oscillated in pitch ±50°at 0.025-0.1Hz.

Results
Double oscillations of BP and HR were induced by both binaural and monaural stimulation in each rat. The double oscillations were commonly induced by stimulus frequencies ≤0.1Hz. Stimulation at higher frequencies commonly induced only single oscillations. Single sinusoids produced increases in BP that slowly declined over several minutes whether give monaurally or binaurally. Double oscillations in BP were similarly induced by tilting rats ±50°at low frequencies.

Conclusion
The facts that sGVS induces both MSNA in humans and VVRs in rats and that single sinusoids induce increases in BP are consistent with the hypothesis that sGVS is channeled through the vestibulo-sympathetic reflex and induces VVRs when appropriately activated. We posit that sGVS sinusoidally activates orientation sensitive central otolith neurons with polarization vectors close to the spatial vertical to produce double oscillations in their firing rates in a manner similar to that induced by pitch and roll tilt. Such otolith neurons have low frequency characteristics. When coupled with the low frequency characteristics of the cardiovascular/baroreflex rhythm, these neurons could drive the large amplitude, low frequency oscillations in BP known as Mayer waves that are commonly associated with VVRs.

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Fluctuating Hyper-G Acceleration Induces Body Temperature Drop in Normal but Not Circling Mice

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Background
Balance disorders are increasingly prevalent and may be due to aging or drug treatments. Diagnosis and treatment of these symptoms is complicated because they may stem from neuromuscular disorders rather than a specific vestibular lesion. The potential to learn compensatory behaviors also interferes with evaluation and treatment, in the clinic and in the lab. Recent studies showing vestibular influence of autonomic functions suggests a new avenue of diagnosing vestibular organ defects: measurement of short-term body temperature changes after controlled fluctuations of centrifugal acceleration. Here, we present a device for producing varying accelerations and show it induces short-term depression of body temperature in intact wild-type mice, but not in mutants or chemically lesioned mice with overt vestibular phenotypes.

Methods
Mice were placed in a 50 ml conical tube and accelerated by a rotating arm, with the tube and mouse held perpendicular to the plane of rotation. An automated controller accelerated the 25.4 cm arm to 1.4 cps, then alternated arm speed between 1.4 and 2.4 cps (2 G and 6 G), maintaining each speed for 10 seconds. After a preset number of cycles, the arm slowly decelerated and the mouse was removed from the tube. Core body temperature was measured just prior to acceleration, immediately after, and at specified subsequent times. Mutant mice (Pou4f3, Grxcr1) with circling and head bobbing, associated wild-types, and wild-types lesioned bilaterally with streptomycin were tested.

Results
Wild-types of all strains exhibited a 2-3°C drop in body temperature within 10 minutes of stimulation, and all returned to baseline within 30 minutes. Most mutant mice exhibited no temperature change; none decreased >0.5°C. Streptomycin-lesioned mice also showed little or no drop in temperature.

Conclusion
Brief exposure to controlled fluctuation of hyper-G acceleration produces a repeatable and substantial temporary drop of body temperature in mice with normal vestibular organs. In contrast, mice with vestibular phenotypes due to mutations or ototoxic lesions show no such temperature change. This acceleration protocol is a safe, non-invasive method of testing for peripheral vestibular dysfunction, and free of errors due to the subject ‘learning’ the testing protocol. These characteristics allow this technique to be used to diagnose naïve subjects and those with less overt symptoms, and also to assess the efficacy of therapeutic regimens.

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Summation of Vestibular, Proprioceptive, and Visual Signals in the Parieto-Insular Vestibular Cortex

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Background
Vestibular signals are pervasive throughout the central nervous system, including the cortex, where they likely play different roles than they do in the better studied brainstem. Little is known about the function of the parieto-insular vestibular cortex (PIVC). This brain region is not well defined by anatomical boundaries, but is the largest cortical region with physiological responses to vestibular stimulation and projects to the rostral vestibular and y-group nuclei.

Methods
Single unit neural activity was recorded in the PIVC of 2 rhesus macaques during combinations of rotation in the horizontal plane at 0.2 Hz of the head (vestibular), body (twisting of body relative to head to stimulate the neck), and visual targets. The head was controlled by a head-fixed motor independent from the rotator moving the primate chair. Both passive and active head movements were utilized. The animals were trained to follow visual targets with either their eye or their head. In this way, complex combinations of visual, vestibular, and proprioceptive stimuli were applied while recording.

Results
We report on the responses of 75 PIVC neurons. The activity of these PIVC neurons correlated with the rotation of the head-in-space (vestibular), the twist of the neck (proprioceptive), and/or the motion of visual targets. Visual target sensitivity was not associated with eye movement, but with target motion itself. Most neurons responded to more than one stimulus. The pattern of visual, vestibular and somatic sensitivities on PIVC neurons displayed a continuous range with some cells more strongly responding to one or two of the stimulus modalities while other cells responded to any type of motion with relative equivalence. The responses to combinations of stimuli were often roughly predicted by summing the expected responses to the component head, neck, and target movements. However, this was not always the case. Responses to vestibular and neck rotation stimuli were almost always similar whether motion was passively applied or the result of a volitional movement of the head.

Conclusion
We found that the PIVC neurons respond to convergent, multisensory inputs related to movement of the animal and external visual objects. PIVC neurons may be involved in keeping track of the motion of external objects and the animal's own movements. This comparison of self and external movement in the PIVC is consistent with insular cortex functions related to monitoring, and explains many disparate findings of previous studies.
The influence of tilt and inter-stimulus interval on the roll aftereffect
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Background
Exposure to a stimulus often biases subsequent stimulus perception in the opposite direction. As a result of this aftereffect a neutral stimulus may be perceived as opposite the adapter. One such aftereffect is the post-roll or "Gillingham" illusion in which after making an abrupt roll pilots erroneously perceive a roll in the opposite direction. The potential influence of gravity in the roll aftereffect has not previously been examined.

Methods
Stimuli were delivered using a 6-degree-of-freedom motion platform. In three trial blocks the test stimulus alone was tested at starting tilts of 9° right/left and upright. The maximum test stimulus amplitude was 3° over 0.5 s (peak velocity 12°/s, peak acceleration 75°/s/s). A two-interval procedure was used to measure aftereffects. The adapting (first interval) stimulus was always 9° of roll over 1.5 s (peak velocity 12°/s, peak acceleration 25°/s/s) with right and left interleaved. Blocks of trials included ISI of 0.5, 3, or 6s. For each ISI, subjects started upright (SU), thus the test stimulus occurred when tilted. In other blocks subjects started at ±9° such that the test stimulus occurred when upright-test upright (TU). Eight human subjects participated (42±18 years). A cumulative Gaussian function was determined from each subject's responses.

Results
The baseline bias was near zero in every subject and in all cases < 0.8°/s. The starting position had no significant influence on the bias(p=0.49). The addition of an adapting stimulus often produced an aftereffect. This aftereffect was observed in 4/8 subjects with the SU at an ISI of 0.5s, in all subjects at an ISI of 3, and 7/8 subjects at an ISI of 6. For the TU condition the aftereffects were less consistently observed. The size of the roll aftereffect was similar between TU and SU conditions. At 0.5 ISI the difference between leftward and rightward adapters was 3.0±2.7°/s (TU and SU similar). The aftereffect was smaller in TU relative to SU at ISIs of 3s (p=0.04) and 6s (p=0.01). This is consistent with aftereffect persisting longer when the subject remains tilted after delivery of the adapting stimulus.

Conclusion
The roll aftereffect causes a perceptual bias even after modest low amplitude roll and persist for long durations when subject remains tilted.
Perceived tilt and translation during periodic versus non-predictable motion
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Background
The ability to accurately separate out the effects of tilt relative to gravity from those of translation is essential to maintaining accurate spatial orientation awareness. Previous studies have often characterized motion perception with periodic oscillations or non-predictable steps and pseudorandom stimuli. The purpose of this study was to compare motion perception during periodic sinusoidal motion (0.11 Hz, ±5°), non-predictable sum-of-sinusoidal motion (0.02 – 0.17 Hz, <±7°) and non-predictable random step motion (<±7°).

Methods
Thirteen subjects were either tilted in the pitch plane about their interaural axis or translated in the fore-aft direction along a 4 m air-bearing linear track using an equivalent acceleration as the tilt stimuli. Subjects were trained to use a joystick to continuously record both the magnitude and direction of perceived motion using a visual display of actual motion during step stimuli of various amplitudes. Subjects then reported perceived position of either tilt or translation in darkness with eyes closed.

Results
During tilt motion, phase errors were smallest during sinusoidal motion and increased during both random step and sum-of-sinusoid stimuli. During translation motion, phase errors were smallest during random step motion and increased during both sinusoidal and sum-of-sinusoid stimuli. Joystick responses were more variable during translation motion, with subjects often losing awareness of their position along the linear track during translation trials.

Conclusion
Our results are consistent with perception becoming predictive during tilt motion that can utilize the gravitational vector as a corrective anchor. The absence of a similar reference during translation in darkness without tilting increases the perceptual uncertainty during either periodic or non-predictable oscillations. We hypothesize that during natural lower frequency stimuli, phase-linking of translation movements with tilt motion and non-vestibular references is important to construct a perception of one's overall motion path.

Assessing Misperception of Rotation in BPPV
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Background
Perception of illusory motion is often the most disturbing symptom of a vestibular disorder. However, these misperceptions of motion are difficult to measure, even though they may be a useful indicator of vestibular function. A
classic example of illusory motion is the vertigo of benign paroxysmal positional vertigo (BPPV), in which perceived rotation arises especially during the Dix-Hallpike maneuver. Therefore, BPPV provides a good example in which to study patients’ perceptions of illusory motion. We introduce a method of using cartoon videos to assist in subjects’ reporting of misperceived motion, and investigate whether or how the reported misperceptions of motion vary between subjects.

Methods
Patients were identified with unilateral BPPV of the posterior canal only. The patients were instructed to pay attention to any perceived rotation while undergoing the Dix-Hallpike maneuver. After the Dix-Hallpike maneuver, the patients indicated the direction of perceived rotation by choosing between cartoon videos of possible perceptions. The videos showed a subject in the Dix-Hallpike supine position, and had stripes on the head that rotated in a certain direction in each video. Each patient was instructed to choose one of more videos to indicate the rotation perceived during testing.

Results
Most patients chose the video with backward rotation about the posterior canal axis of the affected ear, but the majority also chose at least one additional video with rotation about an orthogonal axis. The most-selected videos were backward rotation about the posterior canal axis, ipsilateral rotation about an axis aligned with the body (yaw relative to the body), and ipsilateral rotation about an earth-vertical axis (roll relative to the body). Every combination of one, two, or three of these motions was selected by at least one patient. Nevertheless, the results showed test-retest reliability within subjects.

Conclusion
Patients with posterior canal BPPV report perception of rotation that is not necessarily aligned with the affected posterior canal axis. In addition, the direction of rotation differs between subjects. These results show that perception and eye movements must be studied independently in order to gain a full understanding of vestibular disorders.

554 Insights into prestin function from knock-out/knock-in mice
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The discovery of outer hair cell (OHC) somatic electromotility was followed by a flurry of experiments designed to document this voltage-dependent behavior and to evaluate its role in cochlear amplification. The subsequent discovery of the OHC motor protein, prestin, resulted in the development of knockout mice lacking prestin and showing threshold shift and lack of tuning. In order to obtain a better model to test prestin’s functional significance, knockin (KI) mice were developed to overcome the reduced stiffness and length of OHCs in prestin knockout (KO) mice. For C1 KIs (K233Q, K235Q, R236Q; Oliver et al., 2001; Gao et al., 2007), the peak of the nonlinear capacitance (NLC) function was shifted in the hyperpolarizing direction but the mice showed no change in phenotype. In contrast, 499 KIs (V499G/Y501H; Dallos et al., 2008) showed threshold shift and no frequency selectivity due to the fact that NLC and electromotility were vastly reduced at negative potentials. It was also anticipated that prestin KI mice would not suffer the premature OHC death associated with removal of prestin protein. Although the C1 KI met this expectation, the 499 KI mouse demonstrated an aggressive OHC death even before weaning. Other approaches included development of the chimeric mouse in which OHCs lacking prestin interleaved with normal OHCs to produce a mosaic and a hypomorph where various amounts of prestin protein are expressed in a given cochlea. Experimental results using these mouse models raise challenging questions about the processes by which sensitivity and frequency selectivity are established in the peripheral auditory system. Even with a thorough characterization of each prestin mutation, it may be difficult to extrapolate from in vitro results on isolated OHCs or on cell lines to in vivo performance when the OHC is housed within its normal organ of Corti environment. (Work supported by NIDCD Grants DC00089 and DC010633, and The Knowles Hearing Center).

555 Voltage-Dependent Hair Bundle Motion of Chicken Auditory Hair Cells
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Active hair bundle motion of auditory hair cells has been proposed to be the driving force of cochlear amplification in chickens but there is little direct evidence for this notion. The aim of the experiments was to search for active bundle movements in chicken hair cells and investigate their origin. We made patch clamp recordings, mainly from short hair cells (SHCs), in isolated chicken basilar papilla (E18 – P0; 33 degrees C), and monitored hair bundle motion with dual-phododiode imaging. Hair bundle deflections evoked by a fluid jet generated a mechanotransducer (MT) current with maximum amplitude increasing from 0.3 to 1.6 nA from apex to base of the papilla. The MT current encoded hair bundle
displacements with a 10 to 90 percent working range of 30 to 60 nm. The current was blocked by dihydrostreptomycin or FM1-43 but was unaffected by 10 mM Na-salicylate. Depolarizing voltage steps induced a negative displacement of the hair bundle (away from the kinocilium) of 10 to 40 nm, sometimes followed by a positive overshoot after the voltage step. The voltage-evoked bundle motion consisted of two components of opposite polarity. Blocking the MT current with FM1-43 enhanced the negative motion indicating a positive component linked to gating of MT channels, whereas 10 mM salicylate reversibly blocked the negative movement. A non-linear capacitance change could also be measured in SHCs, also blocked by salicylate, with a peak voltage around +6 mV. Similar results were obtained in tall hair cells. Depolarizing one SHC in preparations lacking a tectorial membrane (TM) induced a displacement in bundles of adjacent hair cells, suggesting a movement of the cell body or cuticular plate. With the TM still present, its radial motion, assayed by imaging attached silica beads, could be produced by extracellular current stimulation. This TM motion was also blocked by salicylate and was of similar size (up to 20 nm) to evoked movements of hair bundles beneath the beads. Our results reveal a novel movement in chicken hair cells which might originate from their cell bodies instead of hair bundles. Movements of the same size for the TM and hair bundle beneath indicate that the active bundle force can be transmitted via the TM to the tall hair cells. The voltage-dependence, non-linear capacitance and sensitivity to salicylate suggest the involvement of chicken prestin in active hair bundle motion.

New Insights into the Structure-Function Relationship of Prestin
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Prestin is a member of the family of solute carrier 26 (SLC26A) anion transporter proteins. It is the motor protein of the cochlear outer hair cells and uniquely functions as a direct voltage-to-force transducer. Although intracellular anions are known to be critical for voltage sensing and conformational change of prestin, the structure-function relationships of prestin remain to be elucidated. In this presentation, we will review recent work from comparative, evolutionary studies using site-mutagenesis, domain swapping and voltage-clamp techniques. On the basis of our new studies using a combination of abinitio structure prediction, 3D folding recognition by threading, homology modeling, molecular dynamics simulations and site-mutagenesis, we propose a new mechanism of how prestin interacts with intracellular anions to generate gating current. (Supported by NIH grant R01 DC 004696 from the NIDCD).

Prestin Structural Biology – Evidence for Oligomerization and Co-operativity
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Since the identification of prestin as the motor protein of the outer hair cell, much progress has been made in elucidating its structure and mechanism, although much more remains to be understood. Recent attention has focused on its oligomeric nature and the possibility that the subunits work cooperatively. The presentation will review the contradictory nature of the evidence and will attempt a consensus view.
The Structural Basis of Electromotility and Anion Transport by Prestin and SLC26 Anion Transporters

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Despite the substantial efforts since the identification of prestin (SLC25A5), the molecular mechanisms that generate its electromechanical ('piezoelectric') activity have remained elusive. Similarly, little is known about the mechanisms that underlie anion transport by the close homologs within the SLC26 anion transporter family. Progress in understanding the molecular basis of SLC26 function has been hampered mainly by the lack of structural information on SLC26 proteins.

We have previously found that non-mammalian orthologs of prestin are electrogenic anion transporters. Generation of chimeras between electromotile prestin and its transporting non-mammalian orthologs enabled us to identify two domains within the transmembrane core region that cooperate in the generation of electromotile activity. This finding defined functional domains of prestin for the first time and demonstrated an overlap in the protein structures that mediate piezoelectric activity in prestin and anion translocation in SLC26 transporters. Moreover, the results pointed to mechanistic similarities between electromotility and transport. However, how both domains function in the generation of electromotility and anion transport remained unknown.

Here we present a 3D structural homology model of prestin. Using a bioinformatic approach to search protein databases we identified a recent crystal structure as the appropriate template for homology modeling. Molecular dynamics simulations showed that the obtained structure is energetically favorable and stable, strongly supporting the suitability of the template used for modeling and confirming the quality of the derived structure. Interestingly, the transmembrane architecture of the model differs from previously suggested topologies, and thus allowed for an experimental test of its validity. We therefore performed a substituted cysteine accessibility scan to determine the location of all predicted TM-TM linkers. The accessibility data are entirely consistent with the structural model but disagree with previously proposed 10 or 12 TM topologies.

The proposed model further pinpoints a central substrate binding site. Point mutations at this site affected anion selectivity both in prestin and its transporting orthologs and strongly altered the efficacy of the competitive inhibitor, salicylate. We conclude that the binding site that confers the anion dependence of electromotility corresponds to the central substrate binding site for anion translocation in other SLC26 transporters.

In conclusion, we present an experimentally validated structural model of prestin that will be highly useful for addressing the molecular mechanisms underlying electromotility and anion transport.

Prestin the atypical Slc26 transporter

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Prestin is a member of the SLC26 solute carrier family and functions as motor protein in cochlear outer hair cells. Whereas other SLC26 homologs transport as anion exchangers or channels a wide variety of anions, prestin is the only member of this family with piezoelectric properties. For many years, it was unclear whether mammalian prestin can mediate anion transport. The pseudohalide thiocyanate (SCN⁻) has been shown to uncouple many anion exchangers, and we thus used SCN⁻ to study whether prestin can also assume “slippage” modes of anion conduction. Mammalian cells heterologously expressing rat prestin exhibit SCN⁻ currents above background levels. The amplitudes of these anion currents are proportional to the number of prestin molecules per cell. Variation of the SCN⁻ concentration resulted in changes of the current reversal potential that obey the Nernst equation indicating that SCN⁻ transport is not stoichiometrically coupled to other anions. Application of external SCN⁻ causes large increases of anion currents, but only minor changes in non-linear charge movements. We conclude that prestin can assume an anion transport mode that is promoted by SCN⁻. However, only a very small percentage of prestin molecules functions as SCN⁻ transporter under these conditions whereas the majority still functions as motor protein. Unitary current amplitudes are below the resolution limit of noise analysis and thus much smaller than expected for pore-mediated anion transport. We conclude that mammalian prestin is capable of mediating electrogenic anion transport and suggest that SLC26 proteins converting membrane voltage oscillations into conformational changes and those functioning as channels or transporters share certain transport capabilities.
The relevance of basic research on perceptual learning to clinical auditory rehabilitation
Beverly Wright

Human adults can improve their auditory perceptual skills through practice, making perceptual training a promising tool to treat communication disorders. Our premise is that a greater understanding of the basic kinetics and mechanisms of skill learning in humans will lead to more effective training strategies. A general model holds that learning occurs over two phases, acquisition and consolidation. The acquisition phase is the actual period of training. The consolidation phase is the period after training ends during which the memories that were formed during acquisition are transformed from a fragile to a more stable state. The consolidation period can last for hours to days and behavioral improvements are often revealed only after consolidation has occurred. Thus, it is thought that the hour during which you are taking a tennis lesson is the period during which you are acquiring the experiences necessary to improve your serve, but it is during the hours after that lesson, when you are engaged in other activities including sleep, that those experiences are consolidated in long-term memory, yielding lasting improvements in performance. We have been addressing these principles in auditory learning by characterizing the circumstances that are necessary for acquisition to occur and by determining the vulnerability of the learning process to intervening events during the acquisition and consolidation phases. For example, we have evidence that (1) for improvement to occur across days requires a sufficient amount of training per day, and that additional daily training can be superfluous, (2) a combination of practice with relevant auditory stimuli and additional stimulus exposures without practice can enhance learning, and (3) the same intervening event (practice on a non-target condition) can have opposite effects depending on whether it occurs during acquisition or consolidation. We also have begun investigating how these characteristics change with age and are affected by sensory and cognitive disorders. For instance, we have identified a training regimen that is effective for adults, but not for most adolescents nor for most adults with dyslexia. Conclusions drawn from learning on fine-grained discrimination tasks have held for speech learning, suggesting that common principles are at play in both domains. Application of these principles could improve clinical training strategies. Further, an individual’s response to perceptual training could be used as an objective, clinical measure to guide diagnosis and treatment of a cognitive disorder. [Work supported, in part, by NIH/NIDCD RO1-DC04453]
Auditory training: What the Brain Gets versus What it Does with it

Arthur Boothroyd
San Diego State University

Background
What the brain gets is auditory (and/or visual) sensory evidence about spoken language plus information from the immediate physical, social, and language context. What the brain does with this evidence is to determine its most likely source. The listener's knowledge (cognitive, social-cognitive, and linguistic) determines both the possible decisions and the value of the context. The listener's skill determines success - defined, here, as making decisions at high speed (set by the talker) while maintaining an acceptably low probability of error. As in reading, speed is optimized by taking maximum advantage of context and using only enough sensory evidence to confirm predictions "beyond a reasonable doubt". For this reason, effective communication is possible in spite of stimulus degradation by noise, distance, reverberation, and poor or deviant articulation. An acquired hearing loss further degrades sensory evidence and reduces the range of conditions over which effective communication is possible, with serious implications for quality of life. A first, and essential, step in addressing hearing loss is to optimize auditory function with hearing aids and/or cochlear implants...in other words, to increase what the brain gets. To deal with the inevitable residual sensory deficit, the listener must adapt to changed and/or diminished sensory evidence by modifying knowledge and skills. Formal auditory training may speed and optimize this adaptation...in other words, it may help improve what the brain does with what it gets. There is evidence of benefit from auditory training but it is not clear: a) what aspects of knowledge and skill change, b) which aspects of training are responsible, c) how listener characteristics affect answers to these two questions, d) whether improvements affect quality of life.

Methods
San Diego State University student Stephanie Bigler, provided three experienced hearing aid users with 5 to 10 hours of story-level training in noise, using a software package developed by the author under a grant to Gallaudet University. Carry-over was measured using a talker and materials not involved in training.

Results
Improvements were found in the recognition of monosyllables in isolation and words in sentences but these were small. The improvements were not likely to be of practical significance...a conclusion supported by self-report.

Conclusions
These experienced hearing aid users may already have be performing as well as they could. Further research will include new aid and implant users. [supported in part by NIDRR H133E080006].

The application of hearing aid technology to cognitive rehabilitation

Brent Edwards
Starkey Hearing Technologies

Hearing aid development has focused primarily on audibility and signal-to-noise ratio improvements for the vast history of its existence. The impact of hearing aid technology on higher-level processing has largely been ignored in the development, prescription, and assessment of hearing aids. While the impact of top-down processing on auditory function has been well documented, only recently has the impact of cognition on hearing aid benefit been demonstrated. Similarly, the impact of hearing loss and hearing aids has only recently been demonstrated to have an effect on cognitive function. Our laboratories, in collaboration with several partners, have investigated the impact that hearing aids have on higher level function. This research includes the development of new higher-level focused outcome measures, demonstrations of the impact of hearing aids on listening effort, measures of a hearing aid’s effect on auditory scene analysis, and diagnostic correlates with cognitive benefit from hearing aids. This talk will review our laboratories’ research projects in these areas, including the motivation for each, and will speculate on the impact of this research on the future of the hearing aid field.

Training the Older Brain to Understand Speech in Noise

Larry Humes
Indiana University

The following facts have been established regarding age-related hearing loss and speech communication. Approximately 40% of older adults have a peripheral hearing loss sufficient to cause speech-understanding difficulties. For those who seek help, typically 10 years have passed from the time a hearing loss is first suspected to the onset of intervention. The most frequent complaint of older adults is that they can hear speech, but can’t understand it, especially in backgrounds of noise. Despite significant technological advances in recent years, contemporary hearing aids still fall short in fully addressing the speech-in-noise difficulties for many older adults. Hearing aid use alone (also known as “acclimatization”)
does not appear to offer significant improvements in speech-in-noise performance through extended periods of use (at least out to 3 years of hearing-aid use).

In addition to these “auditory facts” regarding aging and speech communication, some additional and potentially relevant facts have been established regarding age-related changes in cognition and linguistic processing. In general, in normal healthy aging and for process-related cognitive measures, cognitive function declines steadily for the average adult beginning at about an age of 30 years. On the other hand, product-related cognitive function, such as vocabulary, verbal comprehension and use of context, increases throughout much of adulthood, only declining at or beyond an age of about 75 years.

With this background in mind, we have been investigating an auditory-training regimen designed specifically to improve the ability of older adults to understand amplified speech in noise. The focus of this presentation, after reviewing briefly the evidence supporting the facts outlined above, will be on the conceptual framework underlying our auditory-training regimen and the evidence gathered to date in support of its efficacy. (Work supported, in part, by NIA R01 AG008293 and NIDCD R01 DC010135.)

**Outcome measures in children fit with hearing aids and cochlear implants**

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**Background:** Speech recognition measures are the primary outcome used to validate the effects of sensory devices for children with hearing loss. Clinicians use these measures to determine candidacy (i.e. cochlear implant [CI] vs. hearing aid [HA]), to document progress in speech, language and auditory development and to assess the effects of device parameter selection. As technology improves and CI candidacy expands to include children with more residual hearing, there is a need to incorporate a variety of measures that better reflect real-life listening environments and have applications for clinical practice. Current clinical practice includes fitting and evaluating children who have different device configurations, for example CI and a HA (i.e., bimodal). The fitting of both the HA and CI are critical variables that contribute to outcomes and specifically bimodal benefit. We present data from our lab where several speech recognition measures were incorporated to evaluate the effects of HA frequency response (FR) on bimodal benefit in children.

**Method:** Bimodal fittings with three different FRs were compared to the CI alone condition to assess benefit. Three FRs were evaluated: 1) baseline bimodal fitting with a traditional wide band FR, 2) restricted high frequency gain based on residual hearing and real ear gain, and 3) activated frequency compression. A baseline (A-wide band FR) and two experimental (B1-restricted response, B2-frequency compression) FRs were evaluated in an alternating research design (A/B1A2). Fourteen children (ages 7-21 years) were assessed with measures of speech recognition in quiet and noise, talker variability and localization. Subjective questionnaires were completed by parents and children for all conditions and each child was asked which of the FRs they preferred.

**Results:** Group data revealed no significant differences between mean scores for the three FR conditions in the bimodal condition compared to the CI alone condition across most measures. Group analysis using the FR that resulted in the highest individual bimodal scores revealed that the bimodal condition was significantly better than the CI alone condition across all measures. The best bimodal condition varied across individuals and outcome measures, and the preferred FR varied across individuals.

**Conclusions:** Individual results support consideration of both restricted bandwidth and frequency compression, in addition to the traditional wideband frequency response, for HA settings used in bimodal fittings.

**Pediatric assessment and rehabilitation of hearing loss**

Karen Gordon¹

¹The Hospital for Sick Children

Sub-title: Defining and treating hearing loss in children

Hearing loss is one of the most common congenital anomalies and, when left untreated, results in significant deficits in oral communication. Despite clear effects on the immature auditory pathways during infancy and childhood, it is easy to overlook often subtle signs of hearing problems in young children until speech and language development are already delayed. The use of physiological measures of auditory function have revolutionized the diagnosis of hearing impairment in infants, making it possible to determine how much hearing sensitivity has been lost across the range of frequencies important for understanding speech. Clinically useful measures include otoacoustic emissions, which assess outer hair cell function, and evoked potentials, which measure electrical fields of neural activity at discrete areas of the auditory
pathways. Many countries use these tests to screen for hearing loss in newborns. Treatment of permanent sensorineural hearing loss through auditory prostheses can be initiated based on these measures and refined once the child learns to respond to sound. Current research indicates that childhood hearing loss is heterogeneous with multiple associated genetic mutations affecting specific structures in the cochlea and/or auditory pathways. Advances in treatments must thus both account for the particular deficit involved and the degree of abnormal change suffered. It is clear that we can limit degenerative changes to the auditory system by restricting the time children are left without hearing and we continue to work on how best to restore hearing to each child.

Wrapping it up - What do we know and where do we need to go?
Kelly Tremblay

Understanding the effects of sensory experience on the brain is a long-standing theme of research, crossing all modalities, in the field of neuroscience. In the auditory domain, motivation comes from at least two streams of scientific inquiry: 1) defining normal processes associated with auditory learning; including, but not limited to speech, language, and music; and 2) using the proposed models of learning to develop effective ways of (re)habilitating impaired perception. Here we wrap up this special session by reviewing what is known about the importance of access to sound, as well as the various ways humans are able to make use of this sound. In attempt to translate this knowledge into clinical practice, this summary will emphasize gaps in knowledge where basic and applied scientists can work to improve the hearing health of people with communication disorders.

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An Approach to Drug Discovery: Screening for Survival and Neurite Promoting Activity on Spiral Ganglion Neurons

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Background

One key goal in hearing medicine is to develop interventions that will maintain survival and promote neurite regeneration from spiral ganglion neurons. Given the importance of spiral ganglion neurons for normal hearing and for the function of cochlear implants, one would assume that thousands of drugs and chemicals had already been evaluated for survival and neurite promoting activity in the cochlea. This has not happened, however, for a logical reason: Evaluating just one drug or chemical in the ear at multiple concentrations and time points, for effects on hearing and cochlear anatomy is so labor intensive that it is simply not possible to test thousands of drugs or chemicals in hearing loss animal models. Rather than push forward with unique or unknown compounds in the ear, the field has thus far mainly been occupied with factors that were first vetted in other regions of the nervous system. To focus in on interventions aimed at cochlear neurons, a new approach is required to speed up the process and strike a productive balance between discovery and \textit{in vivo} evaluation.

Methods

Our approach uses a well-characterized and quantified primary culture system of dissociated spiral ganglion from newborn mice. From 8 mouse pups, 192 cultures are generated in 384 well plates, tested, immunolabeled, automatically imaged and quantified. Forty four chemicals and 16 controls can be assayed in quadruplicate using the material generated from 8 mouse pups.

Results

In initial studies we demonstrated that the Rho Kinase inhibitor H-1152 stimulates neurite growth in spiral ganglion cultures. We further demonstrated that other Rho Kinase inhibitors, Fasudil and hydroxyfasudil also stimulated neurite growth, albeit at a higher concentration.

Conclusion

As an example of the utility of our approach, we will demonstrate results acquired in our screen of the NCC clinical collection, a library of over 400 known bioactive compounds. (Supported by ONR grant\#N00014-12-1-0173)

Generation of human inner ear prosensory-like cells via epithelial-to-mesenchymal transition

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Background

Mammalian sensory hair cells are vulnerable to numerous insults, such as overstimulation, aging, genetic orders, and ototoxic drugs. Damage to mammalian sensory hair cell is usually irreversible; therefore, degeneration of hair cells generally causes permanent hearing loss, tinnitus, and other inner ear disorders. Generation of hair cell progenitors with the capability of proliferation and differentiation has been suggested as one of the critical steps in hair cell regeneration. It has been documented that mouse inner ear sensory epithelia possess stem/progenitor cells, which show the ability to proliferate and differentiate in vitro. However, it remains undetermined: (a) how do adult human utricular cells respond when dissociated and cultured in vitro; and (b) whether stem/progenitor cells exist in adult human inner ear sensory epithelia.

Methods

We collected discarded utricles from translabyrinthine surgery and isolated human utricular sensory epithelial cells (HUCs) to explore whether HUCs can proliferate and obtain features of stem/progenitor cells in vitro using reverse transcription polymerase chain reaction (RT-PCR) and immunofluorescence.

Results

When cultured in vitro, dissociated HUCs started to expand on the substrates and express genes and proteins that are widely shown in mesenchymal cells, such as ZEB1, CDH2 (N-Cadherin), FN1 (fibronectin), SNAI2, and VIM (vimentin).
Furthermore, HUCs expressed genes and proteins that are usually present in prosensory cells and stem cells, including BMP4, EYA1, SIX1, DLX5, HES1, JAG1, CDKN1B (P27kip1), LFNG, NOTCH1, POU5F1 (Oct4), NANOG, SOX2, GFAP, and SSEA4. HUCs were observed to proliferate for up to 25 passages.

**Conclusion**
These results reveal that sensory epithelial cells from the adult human inner ear can undergo EMT to re-enter the cell cycle and express genes and proteins that are usually shown in the prosensory cells, which develop into sensory hair cells during development. The outcomes of this study may open avenues for human sensory hair cell generation, which will provide a novel stem cell-based replacement for inner ear disorders, such as hearing loss, tinnitus, and other inner ear disorders.
Dose response of hair cell damage and regeneration in the neonatal mouse cochlea
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Background
Regeneration of β cells in the pancreas has been well studied and different mechanisms of regeneration occur depending on the amount of initial β cell loss (Nir et al., 2007 J Clin Invest; Thorel et al., 2010 Nature). We have previously shown that the neonatal mouse cochlea has the capacity to regenerate hair cells (HCs) after damage in vivo (Cox et al., 2012 ARO abstract #1066). After ablation of ~80% of neonatal HCs, we observed direct transdifferentiation of supporting cells (SCs) into HCs, as well as a new mechanism of HC regeneration where SCs changed cell fate, expressed early HC markers, and divided. We are now investigating the HC regenerative response that occurs after different amounts of initial HC death.

Methods
We used the HC-specific CreER line, Atoh1-CreERTM, to drive expression of diphtheria toxin, fragment A (DTA) with the ROSA26-loxP-stop-loxP-DTA allele. Tamoxifen injections at postnatal day (P) 0 and P1 achieves ablation of ~80% of HCs, while a single tamoxifen injection at P0 results in ~50% HC loss. We used a second CreER line, Atoh1-CreERT2, with tamoxifen injections at P0/P1 to decrease the amount of HC loss to ~20%. HC regeneration mechanisms were then probed using the Hes5-nlsLacZ allele for fate mapping of SCs and BrdU injections to label cells in S phase.

Results
Both direct transdifferentiation and the new mechanism of HC regeneration were observed with the two smaller amounts of HC damage. We are currently creating a model to decrease the amount of HC loss further.

Conclusion
The neonatal mouse cochlea is able to respond to even a 20% HC loss and maintain the capacity to regenerate HCs using both direct transdifferentiation and mitotic mechanisms.

This work was supported in part by grants from the National Institute of Health (DC006471, DC008800, DC010310, and CA21765 the Office of Naval Research (N000140911014, N000141210191, and N000141210775), and the American Lebanese Syrian Associated Charities (ALSAC) of St. Jude Children’s Research Hospital. J. Zuo is a recipient of The Hartwell Individual Biomedical Research Award.
Lineage tracing reveals supporting cells contributing to hair cell regeneration in the neonatal mouse cochlea in vivo

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Background
Mammalian hair cell (HC) loss which leads to permanent hearing loss is long considered irreversible. The discovery of HC regeneration in the avian cochlea provided impetus to readdress this assumption. Several important studies have shown that neonatal cochlear supporting cells (SC), unlike those from older ages, displayed limited progenitor cell characteristics including the ability to proliferate and differentiate into HC-like cells in vitro. The relative paucity of efficient models for eliminating cochlear HC in vivo in both neonatal and adult mammals has limited analysis of the regenerative capacity of these SCs.

Methods
Here, we used the Pou4f3DTR transgenic mice where the human diphtheria toxin receptor is expressed specifically in HCs. HCs were ablated on postnatal day (P) 1 or 6 with toxin injection and tissues examined 1-9 days later. In parallel, we fate-mapped Lgr5⁺ SCs prior to HC loss using the Cre-loxP approach in Pou4f3DTR;Lgr5creERT2;ROSA26RtdTomato mice.

Results
After diphtheria toxin injection at P1, we observed an initial decrease of myosin7a⁺ HCs between P2-P5 (23% of control at P5) followed by a small but significant increase at P6-P7 (51% at P7) in the apex. This recovery in HC number was transient and most pronounced in the apex, and decreased towards the base. At P7 many myosin7a⁺ HCs expressed the prosensory/SC marker Sox2 (apex: 36±8; middle: 9±4; base: 0±0), when none were observed in undamaged, time-matched controls. To investigate whether SCs contribute to these potentially regenerated HCs, we fate-mapped Lgr5⁺ SCs prior to HC ablation in Pou4f3DTR;Lgr5creERT2;ROSA26RtdTomato mice after toxin injection at P1. Compared to controls (Lgr5creERT2;ROSA26RtdTomato), HC loss significantly increased traced tdTomato⁺/myosin7a⁺ cells at P7 (99±5, 23±7, 2±1, respectively), many of which also expressed Sox2 (19±6, 3±2, 0±0, respectively). HC damage induced at P6 led to a progressive HC loss without any regeneration. At P9 when cochleae have lost 20-40% of HCs, we did not detect any myosin7a⁺/Sox2⁺ cells or increase of tdTomato⁺/myosin7a⁺ cells in comparison to controls.

Conclusion
During the early postnatal period, mice have a limited capacity to replenish lost cochlear hair cells in vivo. Regeneration is most evident in the apex and diminishes toward the base. Lineage tracing experiments showed that Lgr5⁺ SCs contribute to the pool of replacement HCs. Ongoing work will characterize the functionality of fate-mapped, “regenerated” HCs in older cochlea.
Ablation of p27kip1 in mouse auditory hair cells results in cell proliferation and long term survival of postnatally produced hair cells.
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Background
p27kip1 has a well-established role in the quiescence of supporting cells within the mouse cochlea. However, whether or not p27kip1 is expressed, or plays a similar role, in hair cells (HCs) remains unclear.

Methods
To test the role of p27kip1 in postnatal HC quiescence, a tamoxifen inducible Cre recombinase transgene driven by a HC-specific Atoh1 enhancer element (Atoh1-CreERTM) was bred into mice homozygous for a p27kip1 floxed allele. As a comparator, the Atoh1-CreERTM transgene was also bred into mice homozygous for an Rb floxed allele. Experimental mice from both lines were induced with tamoxifen at postnatal days zero and one, while control animals received vehicle only. To assess proliferation, animals were injected with the thymidine analogs EdU and/or BrdU and cochlear whole mountings were immunostained for these compounds as well as for the mitosis marker phospho-Histone 3. Hearing function was tested via auditory brainstem responses (ABRs).

Results
Cre mediated disruption of p27kip1 in cochlear HCs resulted in robust proliferation of inner and outer HCs (IHCs and OHCs, respectively), with many cells progressing through, and completing the cell cycle. Interestingly, there were proportionally more EdU positive IHCs than OHCs, and more EdU positive cells were detected in the apical turns of the cochlea than in middle and basal turns. Additionally, following p27kip1 disruption, numerous HCs survived to at least 6 weeks of age, where supernumary HCs were still readily observed in apical turns, and animals retained ABR threshold levels similar to uninduced controls. By comparison, HCs in Atoh1-CreERTM;Rb(floxed) mice also demonstrated robust proliferation, where IHCs appeared more proliferative than OHCs, apical turns were more affected than middle and basal turns, and many cells progressed through the cell cycle. By contrast, however, very few HCs survived to 6 weeks of age after Rb disruption, and these mice exhibited profound hearing loss as measured by ABRs.

Conclusion
Together, the current data suggest that p27kip1 is expressed in cochlear HCs and that it contributes to their quiescence. Furthermore, p27kip1 presents a significant advantage over Rb as a potential therapeutic target for HC regeneration since the disruption of Rb in HCs adversely affects HC survival and hearing function, while ablation of p27kip1 does not.

This work was supported in part by the NIH, the Office of Naval Research, the Hearing Health Foundation, the National Organization for Hearing Research, and the American Lebanese Syrian Associated Charities.

MYC gene delivery to adult mouse utricles stimulates proliferation of postmitotic supporting cells in vitro
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Background
The inner ears of adult humans and other mammals possess a limited capacity for regenerating sensory hair cells, which can lead to permanent sensory deficits. During development and regeneration, undifferentiated supporting cells within inner ear sensory epithelia can self-renew and give rise to new hair cells; however, these otic progenitors become depleted postnatally. Therefore, reprogramming differentiated supporting cells into otic progenitors is a potential strategy for restoring regenerative potential to the ear. Transient expression of the induced pluripotency transcription factors, Oct3/4, Klf4, Sox2, and c-Myc reprograms fibroblasts into neural progenitors under neural-promoting culture conditions, so as a first step, we explored whether ectopic expression of these factors can reverse supporting cell quiescence in whole organ cultures of adult mouse utricles.

Methods
Utricles were dissected from adult Swiss Webster mice, adhered to glass-bottom dishes, and infected with adenoviruses encoding Oct3/4, Klf4, Sox2, the degradation-resistant T58A mutant of c-Myc (c-MycT58A), or GFP under the control of a cytomegalovirus promoter. Following adenovirus washout, utricles were cultured in growth medium (5% FBS). BrdU was included to label cells in S-phase. To promote differentiation, growth medium was replaced with serum-free differentiation
medium containing N2 supplement. The cultures were fixed in 4% paraformaldehyde at the indicated timepoints, labeled with appropriate antibodies, and imaged on a Zeiss LSM 510 microscope.

Results
Co-infection of utricles with adenoviral vectors separately encoding Oct3/4, Klf4, Sox2, or c-MycT58A triggered significant levels of supporting cell S-phase entry as assessed by continuous BrdU labeling. Of the four factors, c-MycT58A alone was both necessary and sufficient for the proliferative response (Fig. 1A). The number of BrdU-labeled cells plateaued between 5-7 days after infection, and then decreased ~60% by 3 weeks, as many cycling cells appeared to enter apoptosis. Switching to differentiation-promoting culture medium at 5 days after ectopic expression of c-MycT58A temporarily attenuated the loss of BrdU-labeled cells and accompanied a very modest but significant expansion of the sensory epithelium. A small number of the proliferating cells in these cultures labeled for the hair cell marker, myosin VIIA (Fig. 1B), suggesting they had begun differentiating towards a hair cell fate.

Conclusion
The results indicate that ectopic expression of c-MycT58A, in combination with methods for promoting cell survival and differentiation, may restore regenerative potential to supporting cells within the adult mammalian inner ear.
A Comparison of the Effects of BDNF and NT3 Gene Therapy on Peripheral Auditory Fibers and Spiral Ganglion Survival in the Deafened Guinea Pig Inner Ear

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Background
Hair cell loss usually leads to secondary regression of peripheral auditory fibers (PAFs) from the basilar membrane area. Spiral ganglion neurons (SGNs) have a variable survival rate in the deafened ear. Cochlear implants (CI) rehabilitate hearing by bypassing the missing hair cells and directly stimulating the remnant auditory neural structures. Treatment strategies to enhance the survival and health of these remnant auditory neural structures may lead to improved CI outcomes. Prior work has demonstrated that BDNF and NT3 enhance SGN survival and induce PAF regrowth following deafening. To determine which of these neurotrophins performs better in bestowing long term SGN survival and inducing PAF regeneration, we compared the effects of BDNF and NT3 on ears deafened systemically or unilaterally.

Methods
Adult guinea pigs were deafened by either local administration of neomycin into the cochlea or systemic administration of kanamycin and furosemide. BDNF and NT3 were delivered to the adult guinea pig inner ear via adeno-associated viral vectors one week following deafening, and all animals were sacrificed three months following treatment. The extent and pattern of PAF regrowth was assessed on whole mount dissections of the cochlea, and SGN survival was assessed on plastic sections through Rosenthal’s canal.

Results
Both BDNF and NT3 promote regrowth of PAFs and enhance survival of SGNs. Ears deafened with neomycin have a complete loss of hair cells and supporting cells in the basal turn, whereas ears systemically deafened have patches of surviving supporting cells. There is no significant difference in the number of PAFs in the basilar membrane area of animals based on the method of deafening; however, in animals systemically deafened there was a negative correlation between the presence of regrown PAFs and the presence of supporting cells.

Conclusion
Gene therapy with either BDNF or NT3 leads to PAF regrowth and enhanced SGN survival, which may in turn lead to improved CI outcomes. Presence of supporting cells may inhibit PAF regeneration into the basilar membrane area.

This work is supported by The Williams Professorship, and NIH/NIDCD grants DC-010412, DC-007634, T32 DC-005356, and P30 DC-05188.
Lithium Alters the Morphology of Neurites Regenerating from Cultured Adult Spiral Ganglion Neurons

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Background
The small-molecule drug lithium (as a monovalent ion) promotes neurite regeneration and functional recovery, is easy to administer, and is approved for human use to treat bipolar disorder. Lithium exerts its neuritogenic effect mainly by inhibiting glycogen synthase kinase 3, a constitutively-active serine/threonine kinase that is regulated by neurotrophin and "wingless-related MMTV integration site" (Wnt) signaling. In spiral ganglion neurons of the cochlea, the effects of lithium or the function of glycogen synthase kinase 3 have not been investigated. We, therefore, tested whether lithium modulates neuritogenesis from adult spiral ganglion neurons.

Methods
Primary cultures of dissociated spiral ganglion neurons from adult mice were exposed to lithium at concentrations between 0 and 12.5 mM. The resulting neurite morphology and growth-cone appearance were measured in detail by using immunofluorescence microscopy and image analysis.

Results
We found that lithium altered the morphology of regenerating neurites and their growth cones in a differential, concentration-dependent fashion. Low concentrations of 0.5 to 2.5 mM (around the half-maximal inhibitory concentration for glycogen synthase kinase 3 and the recommended therapeutic concentration for bipolar disorder) enhanced neurite sprouting and branching. A high concentration of 12 mM, in contrast, slowed elongation. As the lithium concentration rose from low to high, the microtubules became increasingly disarranged and the growth cones more arborized.

Conclusion
Our results demonstrate that lithium selectively stimulates phases of neuritogenesis that are driven by microtubule reorganization. In contrast, most other drugs that have previously been tested on spiral ganglion neurons inhibit neurite outgrowth or affect only elongation. Our results are qualitatively and quantitatively consistent with lithium inhibiting glycogen synthase kinase 3 activity in spiral ganglion neurons. Experiments with additional drugs and molecular-genetic tools will be necessary to confirm that glycogen synthase kinase 3 regulates neurite regeneration from spiral ganglion neurons, possibly by integrating neurotrophin and Wnt signals at the growth cone.

Derailing organ of Corti development: Uncoupling topology from cell type specification reveals causality of signaling

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Neurosensory precursor cells differentiate as hair cells and neurons through progressive restriction of their developmental potential. Proneural basic Helix-loop-Helix transcription factors consolidate this decision to differentiate as neurons (Neurogenin1) or hair cells (Atoh1). Atoh1 has been claimed to be both necessary and sufficient to generate hair cells. However, Atoh1 expression in the brain generates cerebellar or cochlear nucleus neurons, indicating that the ear provides the context for Atoh1 to differentiate hair cells. Even in the ear, Atoh1 expression occurs in cells that do not differentiate as hair cells such as inner pillar cells or sensory neurons. Understanding this complexity requires a better appreciation of intra and intercellular signals in the developing organ of Corti. Current data suggest that three neurosensory precursor types exist in the ear: one pure neuronal, one pure hair cell and a mixed type with the possibility to differentiate as either hair cells or neurons. This switching is apparently mediated by a neurogenic transcription factor, Neurod1. Neurod1 is known to regulate neuronal differentiation, but is also expressed in hair cells. Neurod1 null mutants transform some neurons into hair cells inside the remaining ganglia through de-repression of Atoh1. Likewise, hair cell fate in the apex of the cochlea is influenced by the absence of Neurod1, resulting in premature expression of Atoh1 with a change in hair cell fate: some outer hair cells differentiate to appear more like inner hair cells. Premature and more profound expression of Atoh1 in these mutants can be corrected using haploinsufficiency of Atoh1. This ability of Neurod1 to alter differentiation of hair cells is not possible with Neurog1 knocked into the Atoh1 locus suggesting that at the time Atoh1 is expressed in undifferentiated hair cell precursors these cells are committed to differentiate as hair cells and this fate decision cannot be changed. However, combining 'self-terminating' Atoh1 conditional deletion with knockin replacement of Atoh1 by Neurog1 shows that under these conditions, Neurog1 is able to differentiate hair cells surprisingly normal, indicating that with the exception of an initial early phase for which Atoh1 is essential, other transcription factors might be able to replace Atoh1. Our data suggest that Atoh1 is a differentiation factor needed to drive a specific step in early hair cell development, Hair
cell fate decision happens before that step, Atoh1 expression reflects that decision and other transcription factors can partially replace Atoh1 in later steps.

### Notch signaling in cochlear development: Puzzles and Pathways

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The Notch signaling pathway has been implicated in the development of sensory tissue in the inner ear, and in regulating the proportion of hair cells and supporting cells that develop in each sensory organ. However, the precise contribution of Notch signaling in the development of the cochlea remains unclear. Although lateral inhibitory Notch signaling through the Notch1 receptor can regulate hair cell and supporting cell differentiation, canonical Notch signaling is not necessary for the development of the prosensory domain from which the organ of Corti derives.

We propose a model in which low levels of Notch signaling are required to establish the neural boundary of the organ of Corti through lateral inductive processes, and higher levels of Notch signaling are required to regulate the proportion of hair cells and supporting cells through lateral inhibition. This model is based on the analysis of a series of Notch pathway mutants of varying severity, and on the blockade of Notch signaling with a variety of inhibitory compounds of varying strength and specificity.

### Mechanisms of hair cell regeneration in the zebrafish lateral line system

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**BACKGROUND**

The mechanosensory lateral line system is used by aquatic vertebrates to detect water flow. Hair cells are located in clusters on the surface of the body called neuromasts. Like hair cells of the inner ear, lateral line hair cells are susceptible to damage from ototoxic compounds such as aminoglycoside antibiotics. We are using the zebrafish lateral line system to study mechanisms controlling the regeneration process.

**METHODS**

Zebrafish were treated with the aminoglycoside antibiotic neomycin to kill lateral line hair cells. Transgenic zebrafish were used to monitor hair cell replacement. To identify mutations involved in hair cell regeneration, animals were mutagenized with ethynitrosourea and offspring screened in the F3 generation.

**RESULTS**

After aminoglycoside exposure, regeneration is rapid with complete hair cell renewal seen by 72 hours. Timelapse analysis demonstrates that new hair cells are derived from symmetrically dividing precursors. Regeneration can occur after repeated damage in adult zebrafish with no loss in potential for recovery. Labeling hair cells with a genetically encoded photoconvertible tag demonstrates that hair cells undergo continuous turnover. Label retention reveals distinct regions of the neuromast have different kinetics of turnover. A screen for mutations that alter regeneration has revealed those that block hair cell replacement. Initial development of neuromasts is normal, suggesting that defects are regeneration specific. Identification of genes that regulate regeneration in zebrafish is underway.

**CONCLUSIONS**

Taken together our studies suggest that precursors must be continuously generated to replace hair cells lost to damage or turnover.

### Cell-cell Interactions that Underlie Cochlear Innervation

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The sense of hearing is mediated by precisely wired connections between hair cells and spiral ganglion neurons (SGN) in the cochlea. Hair cells are organized by optimal frequency along the length of the cochlea, with low frequencies in the
apex and high frequencies in the base. Tonotopic information is preserved by the SGNs, which send a single process radially from the ganglion and into the cochlear duct, innervating one inner hair cell. SGNs generally connect with nearby hair cells, only rarely extending processes tangentially within the cochlea. Hence, processes from SGNs in a small sector of the ganglion fasciculate to form radial bundles, thereby creating spokes that radiate along the length of the cochlea. The physical arrangement of SGN processes therefore reflects the logic of how sound information is encoded.

To create the mature cochlear wiring diagram, SGN processes navigate a complex environment that contains a wide variety of cell types. We have been using genetic tools to label and visualize SGNs, efferents, and glia in the developing cochlea. During the earliest stages of differentiation, SGN neurites grow through the mesenchyme to reach the cochlear duct. Once in the cochlear duct, SGN processes encounter support cells and eventually reach the hair cells. In addition, neural crest-derived glia migrate in from the central nervous system and become tightly associated with SGNs and their processes. All of these events occur in parallel for the cochlear efferents, which originate in the brainstem and grow along the eighth nerve and into the cochlea, extending together with SGN processes in the radial bundles. We find that SGN processes are preceded by a wave of glia. Efferents, on the other hand, follow the afferents, spiraling towards the apex upon reaching the cochlea, and then eventually sending processes into the radial bundles and towards the hair cells. To characterize the interactions between SGNs and surrounding cells, we have established a method for live imaging in embryonic cochlear explants. This has allowed us to analyze the pattern of SGN outgrowth and cell-cell interactions in wild-type cochlea and in mutants with abnormal cochlear wiring patterns. By ablating specific cell types and blocking critical signaling pathways, we have begun to elucidate the rules that govern cochlear wiring.

ATP-dependent signaling in supporting cells of the developing cochlea
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Neural circuits that process sensory information exhibit spontaneous, sensory-independent, electrical activity during early development, which has been shown to promote neuronal survival, sculpt electrophysiological properties, and induce refinement of synaptic connections. In the auditory system, neurons fire discrete bursts of action potentials separated by long periods of quiescence prior to hearing onset, events that are initiated by periodic depolarization of hair cells in the developing cochlea. Our studies have focused on understanding the mechanisms responsible for initiating this spontaneous activity, and the effects of this activity on the maturation of cochlea and auditory circuits in the CNS. Studies performed on both acutely isolated and cultured cochleae from prehearing rodents indicate that supporting cells within a developmentally transient expanse of columnar epithelial cells known as Kölliker's organ (also termed the greater epithelial ridge, GER), play a crucial role in triggering hair cell depolarization by periodically releasing ATP. The release of ATP from these cells sets in motion a cascade of events that results in generation of bursts of action potentials in spiral ganglion neurons. Pharmacological and physiological manipulations in excised cochleae indicate that extracellular ATP binds to purinergic autoreceptors (primarily P2Y receptors), causing elevation of intracellular Ca²⁺ and opening Ca²⁺-activated Cl⁻ channels. The resulting efflux of Cl⁻ has two main consequences: (1) it induces cell shrinkage (crenation) due to the concomitant movement of water to maintain osmotic balance, and (2) it forces efflux of K⁺ to maintain ionic balance. K⁺ that is released into the extracellular space accumulates around inner hair cells, inducing depolarization, Ca²⁺ action potentials, glutamate release and eventual excitation of spiral ganglion neurons. In addition, the depolarizing shift in the equilibrium potential for K⁺ that occurs around inner hair cells during ATP-evoked events enhances the release probability of cholinergic efferent fibers. This spontaneous activity declines abruptly after the ear canal is drained of fluid and normal hearing thresholds are established, due to a cessation of ATP release from inner supporting cells. Together, these results suggest that ATP signaling in supporting cells of the cochlea plays a crucial role in initiating spontaneous electrical activity in the auditory system during early development.
Expression of human pendrin rescues hearing in mice lacking mouse pendrin
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Background
Mutations of SLC26A4, the gene coding for pendrin, are a common cause for deafness associated with an enlargement of the vestibular aqueduct (EVA). Slc26a4−/− mice, a model for EVA, develop an abnormal enlargement of the membranous labyrinth and fail to develop hearing and balance. The goal of this study was to determine whether limited expression of human pendrin can prevent enlargement of the membranous labyrinth and rescue hearing and balance.

Methods
A transgene was generated to express human pendrin under the promoter of the B1 subunit of the V-H+-ATPase, Atp6v1b1. This transgene was crossed into Slc26a4−/− mice to generate Tg(+)Slc26a4−/− and Tg(-)Slc26a4−/− as well as Tg(+)Slc26a4+/− and Tg(+)/− mice. Auditory brainstem recordings were used to determine hearing thresholds and RotoRod was used to test balance. RT-PCR was used to determine the expression of Atp6v1b1 mRNA during cochlear development and to differentiate between expression of human and mouse Slc26a4mRNA. Immunocytochemistry was used to evaluate pendrin protein expression without distinction of species origin.

Results
Between embryonic day (E) 14.5 and postnatal day (P) 8 Slc26a4−/− and Slc26a4−/− mice expressed similar levels of Atp6v1b1mRNA in the cochlea. This finding makes it likely that mRNA for human pendrin is expressed at similar levels in Tg(+)Slc26a4−/− and Tg(+)/− mice. Tg(+)Slc26a4−/− mice expressed human Slc26a4 mRNA in the cochlea, however, there was no evidence for significant pendrin protein expression. In particular, there was no evidence for pendrin protein expression in the lateral wall or in the spiral limbus, the site that is known to express Atp6v1b1 mRNA (Karet et al., 1999). Most interestingly, Tg(+)Slc26a4−/− mice developed normal hearing and balance. Hearing thresholds in Tg(+)Slc26a4−/− mice were similar to thresholds in Tg(+)Slc26a4−/− and Tg(+)/− mice and RotaRod tests revealed no difference between Tg(+)Slc26a4−/− and Tg(+)/− mice. In contrast, Tg(−)/− mice failed to develop hearing and balance thereby resembling the established phenotype (Everett et al., 2001; Wangemann et al., 2007).

Conclusions
Limited expression of human pendrin protein in the membranous labyrinth of Tg(+)Slc26a4−/− mice is sufficient to rescue the development of normal hearing. This finding in conjunction with the fact that pendrin expression is required for the development but not for the maintenance of hearing (Choi et al., 2011) opens the prospect that a spatially and temporally limited therapy may be sufficient to restore a life-time of normal hearing in mice and possibly in human patients carrying mutations of SLC26A4.

Introduction
Bryan Pollard1
1Hyperacusis Research Limited

Hyperacusis or “hypersensitivity of hearing” is a challenging disorder that is not well defined, not adequately diagnosed, poorly researched, and with few evidence-based treatment options. As a result, many patients with hyperacusis are unable to find clinicians who can adequately diagnose and present a treatment plan to help with their symptoms.

The prevalence of hyperacusis is difficult to ascertain, as the diagnostic criteria are not standardized. However, a Swiss hyperacusis epidemiology study from 2002 by Anderson, Lindvall, Hursti, and Carlbring found 8 to 9% of postal survey and web responders had some level of hyperacusis (although later investigations found these estimates to be high). This would make the prevalence of hyperacusis higher than other well-researched “rare” disorders (e.g. cystic fibrosis has a prevalence of 0.027%). More epidemiologic research is vital to more accurately determine the number of people who suffer from hyperacusis.

For hyperacusis patients, the impact is far beyond an annoyance, as the world has become increasingly loud. There are many common situations that challenge hyperacusis patients on a daily basis including: social events, noisy work environments, travel, medical diagnostic testing (MRI’s, dental drills), caring for young children, etc.; the list is almost endless. Patients with severe hyperacusis become very limited with what they can do, and it can lead to significant isolation and a reduction in the quality of one’s life.
Hyperacusis refers to abnormally sensitive hearing, such that normally tolerable sounds are perceived as excessively loud and/or as unpleasant. It is not associated with unusually low (good) thresholds for detecting sounds. Rather, hyperacusis is manifested by aversive reactions to sounds that are clearly audible but would not normally lead to such reactions. It is useful to distinguish between cases where sounds are perceived as louder than normal, and cases where sounds are not abnormally loud but are nevertheless aversive.

Greater than normal loudness can occur via several mechanisms: (1) Defective operation of the stapedius reflex can lead to greater loudness of low-frequency sounds. (2) Loudness recruitment associated with cochlear hearing loss involves a more rapid than usual growth of loudness with increase in stimulus level once the elevated detection threshold is exceeded. Although loudness often approaches “normal” values for high input levels, sometimes “over-recruitment” occurs, such that high-level sounds appear louder than normal. This effect can be predicted by a loudness model (Moore and Glasberg 2004). It seems to be a consequence of loss of amplitude compression in the cochlea and reduced frequency selectivity. (3) The gain of the central auditory system may be increased following peripheral hearing loss. (4) The functioning of the efferent system may be impaired. This system serves to regulate the gain of the active mechanism in the cochlea. If a signal is sent from the brain to the cochlea with “instructions” to reduce the gain, but the actual gain reduction is less than it should be, the mismatch may be interpreted as excessive loudness.

In other cases, loudness is not the dominant factor. When fear is the dominant factor, the term phonophobia is often used. Dislike of a sound is described as misophonia. These negative emotions sometimes arise as a result of an unpleasant event or experience occurring at the same time as or associated with sound, but such an event cannot always be identified. Once an association is set up, exposure to the “trigger” sounds can result in an over-activation of the limbic and autonomic nervous systems, causing a “fight or flight” response which is itself unpleasant and reinforces the negative associations with sound.

In conclusion, hyperacusis can occur via a variety of mechanisms, both peripheral and central.

The Impact of Hyperacusis on an Individual’s Life
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The impact of hyperacusis on an individual’s life: Monica Weinberg MD, New York School of Medicine. Dr. Weinberg suffers from hyperacusis

Hyperacusis typically has a significant impact on every part of a person’s life, from their work, their home environment, and virtually every social setting. Depending on the severity of one’s symptoms, hyperacusis patients can develop symptoms of ear pain and worsening tinnitus when exposed to certain sounds greater than ~60-75 dB HL. Hyperacusis patients frequently focus on escaping these pain-inducing sound sources. For some patients, the cycle of pain typically induced from a single new exposure can last from hours to days or weeks. As patients stabilize they are often confused about how to determine the risk level to various social settings. For example how can a person determine if a new cycle of pain may be created from a sporting event or child’s musical performance? Should they wear ear protection to be safe? What can they do if an old filling needs to be drilled out? Can additional damage be done? The anticipation and subsequent impact can be very disabling yet hyperacusis is not classified as a disability, and few in the general population or healthcare professions are aware of hyperacusis.

The impact on the hyperacusis patient’s family and social involvement is also significant as friends see a very different person. The joy that came from seeing a favorite performance or movie is replaced with anxiety and ear pain if the patient risks exposing their ears to the event. There is isolation and loneliness if the patient drops all effort to participate in the key social activities that made up their pre-hyperacusis life. Additional tension and frustration can be added to many relationships unless the friend or family member takes an active role in understanding and supporting the unique challenges the hyperacusis patient now experiences on a regular basis.

It is important for auditory researchers and clinicians to hear from someone who lives with the negative impact of hyperacusis and has found positive ways to cope. Additionally, this presentation will help health care professionals better understand what a patient is experiencing, which will allow them to have a more educated approach in dealing with the patient's symptoms.

The Epidemiology and Etiology of Hyperacusis
David Baguley1
1Cambridge University Hospitals
The epidemiology and etiology of hyperacusis: David Baguley PhD, Audiology, Cambridge University Hospitals, Cambridge UK

Data regarding the epidemiology of hyperacusis is sparse. Indications are that 40% of individuals with clinically-significant tinnitus experience reduced sound tolerance. Regarding the general population, there are few population studies available, but the distinction between dislike of intense sound and decreased sound tolerance in general is not clearly made, and an overestimate of prevalence is possible. Baguley and McFerran (2011) derived an estimate of the prevalence of significant hyperacusis at 2% of the adult population. If correct, this indicates a public health problem of some significance.

There are a number of medical conditions which can involve hyperacusis, including Lyme Disease, Addison’s Disease, head injury, migraine and depression. In many cases, however, the sound tolerance issues remain idiopathic. Associations between Autistic Spectrum Disorders and hyperacusis have been reported, but much further work is required.

This presentation will report what is presently known about the epidemiology and etiology of hyperacusis, and where there are significant gaps in our knowledge. The need for information about the mechanisms of hyperacusis to guide treatment interventions will be discussed.
Diagnosing Hyperacusis and Measuring Hyperacusis Severity
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A key challenge to diagnosing hyperacusis is that we do not have specific physical exam findings or imaging tests that provide a standard to accurately diagnose and assess hyperacusis. Loudness Discomfort Level (LDL) diagnostic testing currently performed by audiologists has a high degree of subjectivity. Additionally, LDL testing can induce anxiety in patients with hyperacusis, as they are required to listen to sounds that push their limits of sound tolerance. There is currently no measure of overall hyperacusis impact that combines both psychological and functional impact. We have developed a brief assessment tool, the Tinnitus and Hearing Survey (THS), which includes questions designed to determine if a patient has any degree of a sound tolerance problem. Using the THS, patients with severe sound intolerance are scheduled for a full assessment using primarily the Sound Tolerance Interview to diagnose the extent of the problem and to determine treatment options. Further testing may include in-clinic trials of ear-level sound generators to determine their potential utility in mitigating the condition. The development of a Hypersensitivity Severity Index would be useful in clinical practice to measure improvement rates of hyperacusis patients. Furthermore, no etiology or underlying mechanism in the auditory pathway has been demonstrated to cause hyperacusis. Possible mechanisms include the following: biochemical changes in the peripheral auditory system, auditory efferent dysfunction, a change in central auditory gain, and alterations in the limbic response systems to loud sounds. This presentation will explore current diagnostic methods and offer suggestions for potential approaches for researchers to explore.

Hyperacusis Treatment Options
William Martin1
1Oregon Health & Science University

Hyperacusis treatment options: William Martin PhD, Oregon Health & Science University.

Currently, primary treatment approaches have centered on sound therapy to desensitize the patients’ responses to certain sounds. A popular approach is a habituation training process based on the Tinnitus Retraining Therapy (TRT) methodology which was established by Jastreboff & Hazel for improving tinnitus patients’ condition. There are numerous approaches that are similar from pink noise for music players to neuromonics. Another aspect is that the etiology of hyperacusis can vary, and therefore it is important to recognize that each hyperacusis patient is different. There may be variability in optimal treatment from patient to patient. This presentation will review the current approaches and the evidence associated with their success.

Comparison Of Hair Cells And Supporting Cells By DeepCAGE Sequencing
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Background
Although supporting cells and hair cells have a common embryologic origin, their phenotypes become fixed in the mature, post-mitotic cochlea, and hair cells do not regenerate in adult mammals. A subset of Sox2-Lgr5-double positive supporting cells can divide and differentiate into hair cells after isolation from neonatal mice and are thought to harbor stem cell potential.

Methods
We isolated inner ear stem cells as neurospheres from neonatal ears and subjected them to differentiating conditions that generate hair cells. We also sorted Atoh1-GFP positive hair cells and Sox2-GFP positive supporting cells from neonatal organ of Corti. We used deepCAGE sequencing in collaboration with the FANTOM5 project to gain further information about expression patterns. Bioinformatics tools such as DAVID and Opossum were used for analysis. Genes related to signaling pathways of hair cell differentiation were confirmed by quantitative RT-PCR analysis of the different cell types.

Results
We compared transcription factor expression in supporting cells and hair cells and assessed inner ear neurospheres for changes in pathway activity and transcription factor expression during differentiation. Several genes expressed in other stem cells in the FANTOM database, HMGA2 and Irx1, were highly expressed in proliferating inner ear stem cells, and several genes expressed after differentiation of inner ear stem cells, such as Twist2 and Hlx, were also higher in hair cells than in supporting cells. Genes such as Foxd3, which maintains progenitor status and is important for stem cell self-
renewal, were expressed more in supporting cells than in hair cells. In addition, pathway imprinting technology and changes in transcripational start sites yielded further insight into differentiation of inner ear progenitors. Wnt, Notch and retinoic acid pathways were active during differentiation. Quantitative RT-PCR analysis of the different cell types confirmed the expression patterns in our bioinformatics screen.

**Conclusion**
Comparison of hair cells and supporting cells indicated progenitor properties of supporting cells, and inner ear stem cells during proliferation used pathways of other progenitor cells. They converted to a more hair cell-like phenotype upon differentiation. A better knowledge and understanding of transcription factor expression in the conversion of inner ear stem cell to hair cell will provide important insights for regeneration of hair cells.

**Investigating Hair Cell Regeneration through Integrated Genomic Datasets**

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**Background**
Lower vertebrates are capable of inner ear hair cell (HC) regeneration, but mammals have little capacity to replace these essential mechano-electrical transducers. We seek to understand the regenerative process by comparing the genetic and regulatory architecture of the regenerating avian utricle and cochlea, and by directly modifying specific pathways during avian regeneration.

**Methods**
We employed next generation nucleic acid sequencing (RNA-seq) to derive complete transcriptome information from regenerating avian utricle and cochlea organotypic cultures. Pure SE were collected at 24 hour intervals over a 168 hour time course following aminoglycoside ablation of HCs. Transcriptomes were computationally assembled from biological replicate samples and were compared to untreated control samples that were cultured in parallel.

**Results**
The utricular and cochlear regeneration time courses share ~2,300 differentially expressed genes. Both datasets contain remarkably dynamic patterns of gene expression that correlate with the onset of the phenotypic conversion of supporting cells (SC) to HCs. There are also major differences in the temporal regulation and specific components of Notch signaling, Hedgehog signaling, and apoptosis pathways between utricle and cochlea regeneration. In particular, Atoh1, Notch, and Hes gene expression profiles and gene family members, differ significantly between the two SE.

The statistically significant up and down regulation of transcription factors (TFs) in the databases—647 and 217 differentially expressed in the cochlea and utricle respectively—suggens complex, and possibly different, regulatory networks are working in the two SE. The overwhelming majority of these TFs have never been investigated in HC regeneration.

Our datasets are a valuable new resource for identifying new HC and SC markers. By statistical clustering methods we have identified a set of ~500 new HC-specific genes (and have validated a subset of these by immunohistochemistry). Several of these are novel HC-specific TFs shared between the two SE.

**Conclusion**
We have derived comprehensive measurements of the transcriptome changes that occur in organotypic cultures during avian utricular and cochlear HC regeneration. The insights gained from these databases suggest that there are a core set of ~2,000 transcriptional changes that overlap between the two SE. These most probably encompass the core signals for HC regeneration. There are also several hundred differences in TF programs between the two SE that may point at pathways for cochlea and utricle SE functional specialization.
High-frequency human middle-ear model with compound eardrum and airway branching in mastoid air cells

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Background
The goal is to construct a high-frequency acoustical/mechanical model of human middle-ear function and validate it against published measurements. The model has novel eardrum and middle-ear cleft elements. The eardrum is represented by two components, one component bounded along the manubrium and the other multi-modal component bounded by the tympanic cavity. The eardrum components are additionally coupled to one another by a time-delayed impedance. The middle-ear cleft is an acoustically coupled system of the tympanic cavity, aditus, antrum, and mastoid air cell system (MACS).

Methods
Modeling of ossicular, cochlear, and oval- and round-window function based on a one-dimensional transfer-matrix approach generally follows past studies. The two eardrum components are coupled by a compliant, lossy impedance. Each middle-ear cleft component is represented by a viscothermal acoustic transmission line. A symmetrical binary airway model with 12 generations of airway branchings is constructed for the MACS, in which airway radius and length decrease to 0.2 mm. The MACS input impedance is efficiently calculated over all airways using recursion. Model parameters are fitted to published measurements in human subject ear canals of energy reflectance (0.25-13 kHz) and eardrum input admittance (0.25-11 kHz), and to published temporal-bone measurements of sound pressure in scala vestibuli and scala tympani (0.1-11 kHz). Intracochlear pressure measurements with middle-ear cavities open are modeled using a radiation impedance as the terminating impedance.

Results
The MACS model has total volume of 6.4 ml and surface area of 107 cm², similar to mean dimensions from CT scan analyses of temporal bones. The middle-ear model optimization produced an adequate fit to all data. The two-component eardrum with time delay helps fit intracochlear pressure responses. A multi-modal representation of the eardrum and the high-frequency transmission-line model of the middle-ear cleft help fit ear-canal responses. The eardrum input reactance is small at high frequencies (i.e., not mass controlled). The high-frequency eardrum resonances and the MACS model are critically important in explaining this, and in predicting the frequency dependence of energy reflectance. The ability to accurately explain intracochlear pressure data also explains the input pressure difference acting on the cochlea.

Conclusion
The model accurately fits middle-ear responses to high frequencies, and is well suited for the study of middle-ear dysfunction and for integration into models of cochlear mechanics.
High-speed Video Analysis of Ossicular Chain of Near-intact Guinea Pig Middle Ear
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Background
The actual, acoustic vibration of ossicular chain in the middle ear has not been easy to analyze.

Methods
We investigated ossicular movement in the intact middle ear in response to acoustic stimulation using a high-speed videocamera and video analysis software. The high-speed videocamera could record analyzable ossicular motion up to 1kHz. We made visual access to the middle ear of deeply anesthetized guinea pig by opening ventral wall of otic capsule without injuring sound conductive structures from external auditory canal to oval window.

Results
The stapes showed reciprocal movement in the same frequency with the stimulating tone, and with amplitude partially proportional to the stimulating sound intensity. Injury to the tympanic membrane or round window influenced the mechanical properties of the stapedial motion. When the middle ear was stimulated with multiple tones, the stapes showed complex waveforms whose original stimulating frequencies could be later analyzed and recovered after Fourier transformation.

Conclusion
Our experimental setup was capable of evaluating the conductive hearing regardless of the status of sensorineural hearing or even life. The video analysis may become a powerful tool to investigate middle ear physiology.
Biomechanical Properties of Human and Porcine Auricular Cartilage

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Background

Genetic defects involving mutations of collagen subtypes often result in phenotypic abnormalities of the external cartilaginous auricle as well as hearing. Several studies have detailed the biomechanical properties for septal, costal, and articular cartilages. However, the biomechanical properties of auricular cartilage are largely undescribed. Therefore, it is of interest to determine the biomechanical properties of the cartilage of the external ear.

Methods

Auricular cartilage specimens were harvested from six fresh porcine and six fresh human cadaveric ears. Mechanic properties were determined with an Alliance RT/30 MTS material testing machine. For porcine cartilage two punch biopsies were taken from each ear at a proximal (adjacent to the mastoid), mid, and distal site along the cartilaginous framework. Duplicate specimens were taken and placed in Lactated Ringers solution. The first test that was run on each porcine punch biopsy specimen was a stress relaxation confined compression test. The parameters for this test included confined compression in a compression chamber 6 mm in diameter, to 10% strain, 500N load cell, a preload hold time of 10 min, test speed of 300 um/sec, and a test temperature of 37 C. After overnight relaxation, they were then subjected to a 60% strain unconfined compression test, also carried out using the 500N load cell. The preload was maintained for 3 minutes duration.

For human cartilage, the ear was first subjected to a whole ear helix down unconfined compression test in order to determine overall flexibility/stiffness of the specimen—a method previously unreported in literature. Afterwards, punch biopsies were taken from each ear and subjected to both 10% strain confined compression test as well as the 60% unconfined compression test as described above.

To assess for regional differences in the complex conformation of the human auricle, biopsies were obtained from standardized locations: root of the helix, posterosuperior helix, triangular fossa, conchal bowl, and tragus.

Results

Aggregate modulus and offset yields are provided for porcine and human auricular cartilage. A quasilinear viscoelastic model from stress relaxation data and nonlinear elastic curve using Kenedi-Fung model from unconfined compression data are proposed with graphic representative fits.

Conclusion

Defining robust auricular cartilage biomechanical data allows for comparative indices in analysis of genetic defects affecting collagen subtypes that may have phenotypic biomechanical variation of the external cartilaginous auricle. Both human and porcine auricular cartilage is characterized.
Simulator and Mechanical Model Development for Middle Ear Surgery
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Background
Otologic surgical simulators currently available do not allow interaction with the ossicular chain. To improve teaching interest of these tools, a simulator based on a finite element mechanical model of the ossicles as well as their ligaments and muscles was developed. The objective of this study was to evaluate the mechanical behavior of this model with sound transfer function and sound pressure (tympanometry) simulations.

Methods
A surface model of the ossicular chain and muscles of the tympanic cavity was created from a three-dimensional anatomical data published. Ligaments and muscles were implemented according to the anatomical and physical characteristics reported in literature with Blender (Amsterdam, Netherlands). The model was integrated into the medical simulation software, SOFA (INRIA, France). Mechanical parameters (Young's modulus, density, Poisson's ratio ...) of the ossicles were defined based on the literature's data. The experimental results of the transfer function and forces pressures were compared to results from studies on temporal bones and other finite element models.

Results
A sine wave was applied to malleus and the vibration of the stapes was measured. At 250 Hz and 90 dB SPL, the peak-to-peak displacment of the stapes footplate was 22.4 nm. It increased slightly to 53.8 nm at 1000 Hz before falling sharply to 0.54 nm to 8000 Hz. These measures were in good agreement with measurements on human temporal bones. The evaluation by simulated tympanometry consisted to apply a force to the umbo corresponding to the variation of the pressure in the external ear meatus and to observe the motion of the umbo. These tests showed displacements similar to temporal bones measurements. In case of simulation of annular ligament ankylosis, the displacement decreased slightly whereas a removal stapes yielded to higher motion for high pressure.

Conclusion
Tests have shown that the mechanical behavior of our model was compatible with the experimental data. The mechanical model combined with our surgery simulator was in accordance with the published data on the temporal bones. This model will be used as a training tool for residents on various surgical pathologies.
Motion of the Tympanic Membrane in Acute Otitis Media Model of Chinchilla Measured with Scanning Laser Vibrometry

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Background
Acute otitis media (AOM) arises from acute infection of the middle ear and is caused in about 70% of cases by bacteria. Haemophilus influenzae, a bacterium found in the human nasopharynx, is an opportunistic pathogen that now is the most common cause of human otitis media. There are no reports describing middle ear biomechanical function changes in AOM induced by this Gram-negative bacterium.

Methods
Sound-induced motion of the tympanic membrane (TM) in eight chinchillas with AOM was measured using a scanning laser vibrometer (PSV-400, Polytec). AOM was produced by transbular injection of H. influenzae strain 86-028NP into the left ear and the right ear served as the uninfected control. Four days after challenge, the auditory brainstem response (ABR) and wideband tympanometry were conducted in the animals. The full view of displacement of the TM was acquired by PSV system when a multi-tone stimulus was applied on the TM at the ear canal from 0.2 to 10 kHz.

Results
The wideband tympanograms show that all AOM ears had negative middle ear pressure, and the energy absorbance in AOM ears was lower than that in control ears. ABR thresholds in AOM ears were increased compared with control ears. This finding indicated that hearing loss was associated with AOM even at this early time post challenge of the chinchilla ear with bacteria. The displacements at the umbo and four quadrants of the TM show that there was a significant reduction in the superior-posterior and inferior-posterior quadrants of the TM at frequency below 2 kHz in AOM ears compared to controls. At higher frequencies (2-10 kHz), the reduction was observed only in the superior-posterior quadrant.

Conclusion
3D motion of the entire TM in H. influenzae-induced AOM ears of chinchillas was characterized using the scanning laser vibrometer with the measurements of ABR and wideband tympanometry. The results suggest that there probably exists a correlation between 3D motion of the TM and sound energy transmission into the middle ear as measured in control and AOM ears. (Supported by NIH R01DC011585)
Department of Defense Hearing Center of Excellence: Providing Facilitation of Translational Research in Auditory Injuries

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Background

Military service members are routinely exposed to hazardous environments leading to auditory injuries which diminish quality of life and affect the war fighters’ ability to hear and communicate safely on the battlefield. The US Department of Veteran’s Affairs has documented that impairment of the auditory system currently affects over 1,500,000 individuals, accounting for the two most prevalent disabilities, tinnitus and hearing loss, claimed among all veterans.

In 2009, Congress established the Department of Defense Hearing Center of Excellence (HCE) to address prevention, diagnosis, mitigation, treatment and rehabilitation of hearing loss and auditory system injury for the DOD and the VA. The Air Force was designated as the Lead Component with the goal of ensuring optimal DOD and VA collaboration.

Methods

The HCE research mission is “to facilitate a DoD- and Veteran-wide practice-based research network which will coordinate the identification of and solutions to unsolved problems related to the prevention, diagnosis, mitigation, rehabilitation, treatment, and restoration of hearing loss and auditory injury among US military and Veteran populations.” Focus areas include:

- Provide research tools that promote, and encourage involvement, and remote partnerships among a collaborative network of clinicians, scientists, and cohorts.
- Longitudinal tracking database to identify injury patterns and best clinical practices.
- Prioritized translational strategies for research outcomes.
- Scientific steering and review of auditory research projects and programs.
- Streamlined administrative processes

Results

The HCE has established the DoD Auditory Research Working Group, and expanded it to include technical experts across the VA and NIH, to include a Scientific Advisory Board capacity. The HCE will work with portfolio managers to identify all Federal hearing and balance projects, as well as to serve as a central coordinator for academic/industry collaboration. The HCE has organized several work streams through this networked effort. The Hearing Center of Excellence has identified potential value in integrating research efforts with sister DoD CoEs and has developed a model and initiated discussion with other CoEs to develop a roadmap based on this common model.

Conclusion

The intent of organizing a transparent, coordinated practice-based research network is to encourage and facilitate the conduct of research, and the development of best practices. Based on the foundation of information management, outreach and allied research, the ultimate measures of HCE success will be the effect of this networked system on the prevention of hearing loss, and the outcomes of the health care delivered to those that suffer auditory injury.

The public health importance of noise-induced hearing loss

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Although cochlear physiologists and anatomists had used excessive noise in their labs as a tool to study the effects of damageto the cochlea or auditory nervous system for decades, the general problem of noise-induced hearing loss as a medical condition that required assessment and rehabilitation awaited the end of World War II, and the resultant need to treat the “casualties” who were returning from combat. In his memoirs, Hallowell Davis detailed the development of this new field, which he and his colleagues coined “Audiology” because the military personnel used the term “Auricular Training”, which sounded like teaching them to wiggle their ears! In the 1950’s and 1960’s the basic science work of Bekesy, Davis, Wever, Lawrence, Kiang, Zwislocki, Schuknecht, Hawkins, Tonndorf, and others provided important, new information about cochlear function, anatomical correlates of noise injury, and behavioral effects of excessive noise exposure. In industry, state worker’s compensation laws were introduced to compensate individuals whose hearing was damaged by occupational noise exposure, and increasing efforts to reduce NIHL in the military were introduced. This combination of events led to the implementation of federal regulations governing exposure to noise in the workplace, first established as the Walsh Healey Act in 1969, and then promulgated by OSHA as the Noise Control Act of 1972. This act
Noise-related hair cell death and preservation of endolymph/perilymph boundaries in the Bohne labyrinth
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Cochlear hair cell injury and loss are the best predictors of noise-induced permanent threshold shifts (NIPTS) in animal models. Hair cell cuticular plates form part of the boundary of the endolymphatic space. Thus, large or prolonged openings (or gaps) in the reticular lamina may lead to a decrease in the endocochlear potential (EP) and elevate potassium levels in the organ of Corti, potentially amplifying injury. Such openings might take the form of ‘holes’ left by degenerating hair cells or reticular lamina ‘ruptures’ caused by direct mechanical trauma. Selective pressure to save what can be saved, and to preserve as much hearing as possible, might be expected to favor efficient closure of holes and ruptures and quick restoration of endolymphatic boundaries. Some causes of hair-cell loss (aging, inflammation) are evolutionarily old, while damaging noise is ‘new’. It is unclear whether noise, aging, inflammation, and ototoxins engage the same processes for disposal of damaged hair cells, or how the mode of cell death (necrotic/apoptotic/3rd pathway) modifies death processes (Bohne et al., HR 2007; Lee et al., Open Otorhinolaryngol. J. 2008; Bohne & Harding, ARO 2012). Few studies have been designed to detect holes or ruptures, or to distinguish the exposure conditions or models that favor one form of injury over another. In chinchillas and guinea pigs, holes may form after even moderate noise exposure (Bohne & Rabbit, HR 1983; Fredelius et al., Acta Otolaryngol. 1988; Ahmad et al., HR 2003), while ruptures require much higher levels (Bohne, Effects of Noise on Hearing 1976; Henderson et al., HR 1994; Wang et al., JARO 2002). I will consider evidence for holes versus ruptures as distinct processes, and the implications for cochlear damage and protection.

You say “Toughening”, I say “Conditioning”, let’s call the whole thing off
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You say toughening and I say conditioning. Toughening, conditioning, toughening, conditioning. Let’s call the whole thing off! Both toughening and conditioning are active processes induced by low-level, non-damaging acoustic stimulation that creates long-term protection to subsequent detrimental forms of noise trauma. This phenomenon is now shown to occur in a variety of mammals, including gerbils, chinchillas, guinea pigs, rabbits, rats, mice and human subjects. Different paradigms have been proven successful in preventing a variety of pathological changes to the auditory system. Even though toughening and conditioning experiments are well-documented methods of protecting against hearing loss, the underlying mechanisms have not been well characterized. Several mechanisms have been suggested to play a role in auditory protection by sound conditioning including the activation of antioxidant enzymes, inhibition of apoptosis, increased efferent activity and glucocorticoid activation. This review will discuss these different mechanisms.

What Otoacoustic Emissions Have Told Us About Noise-Induced Hearing Loss
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Soon after the discovery of otoacoustic emissions (OAEs), early reports described their general properties in normal ears in terms of their spontaneity, or their production with various types of acoustic stimuli. The next phase of articles illustrated the effects of various hearing impairments on the assorted OAEs. This presentation will provide an overview of
prior studies that used OAEs to investigate NIHL. Such investigations utilized research in experimental models, attempts
to detect occupational NIHL, or clinical case-study descriptions of emissions in patients with NIHL symptoms. Given that
the initial pathology underlying NIHL entails damage to, or destruction of, the organ of Corti’s outer hair cells (OHCs), it
was expected that OAEs, which are generated by OHCs, would faithfully describe the behavioral audiogram in humans
and/or the underlying cochlear pathology in experimental models. On a population basis, this expectation was usually
born out. However, in general, using OAE tests, it is difficult to predict either the pattern of behavioral hearing or the
underlying cochlear pathology on an individual basis. Among the reasons for explaining disagreements between the
various response measures is the field’s present knowledge about distinct and complex sources that contribute to the
generation of OAEs. Nevertheless, current research findings will be discussed that hold promise for the ability of future
OAE tests to detect early NIHL, and to accurately predict its associated hearing pattern and/or cochlear pathology.

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Tinnitus and Hyperacusis: Involvement of Auditory and Nonauditory Structures
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Background: Exposure to intense noise or ototoxic drugs not only results in hearing loss, but often leads to tinnitus and
hyperacusis. The aberrant neural activity responsible for tinnitus and hyperacusis was believed to originate within the
classical auditory pathway; however, more recent data suggest the potential involvement of nonauditory structures in the
CNS. Here we review some of the neurophysiological changes seen in the cochlea and CNS when rats are treated with a
dose of salicylate that induces tinnitus and hyperacusis-like behavior.
Methods: Operant conditioning techniques and the startle reflex were used to identify doses of salicylate that produce
tinnitus and hyperacusis-like behavior in rats. Afterwards, we characterized the aberrant patterns of neural activity in
the cochlea, auditory pathway and nonauditory sites.
Results: High-dose salicylate (i.p.) caused a frequency-dependent loss in DPOAE and reductions in CAP amplitude;
losses were greatest at low and high frequencies and least in the mid-frequencies (10-20 kHz). Salicylate induced
threshold shifts in the central auditory pathway that mirrored the cochlear losses, but did not significantly alter
spontaneous activity. Despite the threshold shifts and reduced cochlear output (hypoactivity), sound-evoked responses in
auditory cortex, medial geniculate, amygdala, striatum and hippocampus were larger than normal at suprathreshold
intensities (hyperactivity). Moreover, many low-CF and high-CF neurons in auditory cortex and amygdala shifted their CF
into the mid-frequencies. These up-shifts and down-shifts resulted in an over-representation of mid-CF neurons; the
expanded mid-CF region lies near the tinnitus pitch induced by salicylate. To eliminate peripheral drug effects, salicylate
was applied to the amygdala or auditory cortex, while recording from the auditory cortex. In both cases, salicylate
increased sound-evoked activity in auditory cortex at suprathreshold levels, but did not alter threshold.
Conclusion: Salicylate exerts potent effects on the cochlea, central auditory pathway and non-auditory regions of the
CNS. Salicylate-induced thresholds shifts originate in the cochlea whereas sound-evoked hyperactivity, a possible
correlate of hyperacusis, likely originates at auditory and nonauditory sites in the CNS. The salicylate-induced tonotopic
shifts seen in auditory cortex, which may underlie tinnitus, likely results from both peripheral and central changes.
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Translating Data into Knowledge and Action: Challenges in Evidence-Based Hearing Loss
Prevention
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Background: Evidence-based practices seek to ensure that the best available scientific evidence is used in clinical
decision making. Such approach requires assessing all scientific evidence available on the risks and benefits of
interventions and diagnostic procedures. Evidence quality is assessed based on the source type (from meta-analyses and
systematic reviews of randomized clinical trials as high-quality sources, down to conventional wisdom as low-quality
ones), currency, statistical validity, clinical relevance, and peer-review acceptance. The 2009 American Recovery and
Reinvestment Act included a provision for federal funding to investigate how different interventions compare to each other.
The Act called on the Institute of Medicine to recommend comparative effectiveness research priority topics. Their
recommendations are in the report Initial National Priorities for Comparative Effectiveness Research (http://www.nap.edu/catalog/12648.html). The need for research on hearing loss interventions was placed in the highest
priority group. This recommendation underscores the human and societal costs of the condition. The risk of hearing
impairment increases with age and is exacerbated by exposure to noise, particularly at work. This risk can be minimized
by reducing noise levels to 85 dB(A) or less. Many countries have mandated hearing loss prevention programs when
noise exposures cannot be reduced to this level. However, the continuing high rate of noise-induced hearing loss raises concerns on the effectiveness of these programs.

Methods: Recent Cochrane Reviews investigated various initiatives and mechanisms (e.g., legislation, proper hearing protector usage, etc.) to determine which work best to either promote the use of hearing protections, and/or reduce noise exposure or hearing loss among workers.

Results: Results from intervention effectiveness studies on hearing loss prevention do not provide evidence to support current practices. There is consensus in the literature that some interventions improve the use of hearing protection devices compared to non-intervention; there is low quality evidence that legislation can reduce noise levels in workplaces, and contradictory evidence that prevention programs are effective in the long-term. Most reported interventions focus on the use of hearing protectors, and effectiveness depends on the quality of the implementation of prevention programs. Substantial noise control can be achieved in the workplace, with no evidence of this practice in the literature.

Conclusions: Better and large scale implementation of technical interventions and evaluation of their long-term effects are necessary to identify the most effective strategies for reducing occupational hearing loss.

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Frequency Discrimination through Phase Locking
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Background
Humans can discriminate frequencies that are only about 0.2% apart when the frequencies are below 4 kHz. For comparison, a semitone in Western music represents a frequency step of about 6%. Frequencies higher than 4 kHz are increasingly hard to differentiate; as an example, the frequency-discrimination threshold at 6 kHz is about 1.5%. This contrasts with tuning curves of auditory-nerve fibers, which are considerably sharper for high-frequency than for low-frequency fibers. The enhanced frequency discrimination below 4 kHz has been hypothesized to involve the phase locking of specialized neurons in the auditory brain stem that fire at a preferred phase of stimulation.

Methods
We performed psychoacoustic experiments to directly measure the effect of phase locking on frequency discrimination. We generated tones based on a single frequency but in which a random phase change occurred every few cycles. These tones were presented to subjects who performed a frequency-discrimination task. In a theoretical work we then investigated how the frequency information contained in phase locking might be read out in neural networks. We specifically used numerical and analytical techniques to study a class of random neural networks in which signal propagation from one neuron to another induces a time delay and in which the neurons perform coincidence detection.

Results
Our psychophysical experiments showed that random phase changes systematically impede our ability to discriminate frequencies. The frequency resolution in the presence of phase changes is bad as that for short tones whose duration is limited to the time between two subsequent phase changes. We found that the neural networks can perform sharp frequency discrimination when stimulated with phase-locked input: different frequencies induce different neural activity patterns. The frequency resolution achieved by this means depends on the noise in the phase locking and can, for realistic values, reach the minute frequency differences of around 0.2% that have been measured in humans.

Conclusion
Our psychophysical experiments provide direct evidence that phase locking is employed for frequency discrimination. Our theoretical study on a class of random neural networks with temporal delay and coincidence detection offers a framework for how the brain can read out the temporal information contained in phase locking.
Spectral Ripple Discrimination in Infants: Effect of Ripple Depth and Envelope Phase Randomization

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Background
Spectral ripple discrimination (SRD) measures sensitivity to spectral change across a broad frequency range. In infants, SRD might be affected by developmental differences in non-spectral factors, such as intensity resolution, attention, or listening strategy. In Experiment 1, we measured infants’ and adults’ SRD thresholds at various ripple depths to determine the spectral modulation transfer function (SMTF). It was predicted that SMTF height would be lower in infants due to immature intensity resolution and/or attention, while SMTF slopes would be equivalent between age groups due to mature spectral resolution. In Experiment 2, two stimulus conditions were used to determine whether either age group relied on within channel listening strategies. It was predicted that availability of within-channel intensity cues would not affect SRD in either age group.

Methods
Participants were normal-hearing 7-month-olds and young adults. Stimuli were broad-band noise bursts filtered by a full-wave rectified sinusoidal spectral envelope with logarithmically-spaced peaks. The Observer-based Psychoacoustical Procedure was used to determine the highest ripple density at which a participant could detect a 90\degree phase shift in the spectral envelope. In experiment 1, ripple depth was varied between subjects and SRD threshold in ripple density was assessed. In experiment 2, two stimulus conditions were created based on the starting phase of the spectral envelope: Constant (starting phase was always 0\degree) and Randomized (starting phase varied from 0\degree to 270\degree). Stimulus condition was varied between subjects and SRD threshold assessed.

Results
In experiment 1 the effect of ripple depth and age group on SRD threshold was measured using ANOVA. Main effects of age-group (adults with better thresholds than infants) and ripple depth (better thresholds at higher ripple depths) both reached significance. The interaction between age-group and ripple depth did not reach significance, suggesting that SMTF slope was similar between age groups. Results from Experiment 2 preliminarily suggest no significant main effect of stimulus condition on SRD thresholds in either age group.

Conclusion
As expected, SMTF slopes were similar between infants and adults, reflecting mature spectral resolution by 7 months old. Differences in SMTF height between age groups, possibly reflecting immature intensity resolution, illustrate that SMTF slope is a better measure of spectral resolution in infants than SRD threshold at a single ripple depth. Preliminary data from Experiment 2 suggest that neither infants nor adults rely heavily on within-channel intensity cues in an SRD task.
Comparative Measures of Temporal Coherence
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Background
Being able to hear and isolate communication signals in the environment is important for both humans and animals. In humans, we know that basic acoustic parameters such as frequency, loudness, and timing, from different sound sources are important in the perceptual isolation or streaming of multiple auditory objects (Bregman, 1990). Relative to humans, however, little work has been devoted to how animals isolate sound sources to create auditory objects and even less is known about the comparative psychology of auditory stream segregation. Frequency separation of sounds is arguably the most common parameter studied in auditory scene analysis and has been shown to be an important feature for streaming auditory objects in both humans and birds. However, frequency separation of sounds is not the only factor influencing auditory scene analysis, as the relative timing of sounds in the acoustic environment appears important as well (Elhilali, et al., 2009, van Noorden, 1975). Elhilali et al. (2009) found that in humans, synchronous tones are heard as a single auditory stream, even at large frequency separations, compared to asynchronous tones with the same frequency separations, which are perceived as two sounds. These findings demonstrate how both timing and frequency separation of sounds are important for auditory scene analysis. It is unclear how animals such as budgerigars (Melopsittacus undulatus) would perceive synchronous and asynchronous sounds.

Methods
In the present study, both budgerigars and humans were trained and tested using the same operant conditioning procedures to investigate the perception of synchronous, asynchronous, and partially overlapping pure tones and budgerigar contact calls.

Results
Results illustrate that budgerigars segregate partially overlapping sounds in a manner predicted by computational models of streaming. Additionally, both budgerigars and human listeners have trouble segregating harmonic stimuli compared to non-harmonic stimuli of the same duration. These results support previous research illustrating that harmonic sounds most often originate from the same sound source and are more likely to be perceived as one stream (Carylon, 2004; Alain, 2007). Interestingly, for both budgerigars and humans listeners, overlapping budgerigar contact calls and sine-wave bird calls are more likely to be segregated than pure tone stimuli with the same temporal overlap.

Conclusion
These results further support the idea that frequency separation has little influence in how overlapping complex, communication stimuli are segregated, considering the two sets of stimuli fall within the same spectral range.

Auditory streaming of syllables: words fuse more readily than non-words
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Background
The extent to which auditory streaming is influenced by high-level cognitive processes remains a matter of debate. We investigated whether streaming is influenced by linguistic processes, by studying whether streaming of a repeated syllable depends on whether it is a word or a non-word.

Methods
We presented listeners with single spoken syllables (a real word, "stem", "stone" or a non-word, "stome", "sten") repeated every 673 ms for 150 sec. Listeners continuously judged whether they heard the syllables as fused or whether the initial /s/ of each syllable formed a separate stream. We obtained an objective measure of streaming by asking listeners to detect an occasional silent gap of between 20-50 ms long, inserted after the initial /s/ of some “target” syllables.

Results
After a few seconds the initial /s/ was reported to stream apart from the remainder of the sound and the percept then fluctuated in a bi-stable manner between the veridical syllable and a two-stream percept of an isolated fricative /s/ plus the remainder of the syllable. This is an instance of the “Verbal Transformation Effect” (VTE); the lack of aspiration when a /t/ is preceded by /s/, leads to a streamed percept containing the words (“den”, “dome”) or non-words (“dem”, “dohne”). Note that this streaming effect transforms words (“stem” and “stone”) into non-words (“dem” and “dohne”), and vice-versa (for “sten” and “stome”). Performance on the gap-detection task was better at times when the listener reported hearing a
single stream than when the /s/ was streamed off. Importantly, it was also better when the fused percept corresponded to
a word than when it corresponded to a non-word. This provides evidence that the segregation of the sequence into two
streams, or, conversely, its integration into a single stream, was influenced by the lexical status of the repeated syllables.
Note that this finding cannot be attributed to low-level acoustic-phonetic factors, since the vowel (/o/ or /e/) and final
consonant (/m/ or /n/) occurred equally in lexical and non-lexical stimuli.

Conclusion
Our results are consistent with the following conclusions: a) auditory streaming of speech sounds is affected by lexicality,
b) the verbal transformation effect can be driven by streaming, and c) perceptual organisation of speech is sensitive to
lexical status. These findings illustrate how high-level cognitive processes influence auditory and speech perception.

Rhythm perception in macaque monkeys
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Background
The capacity to rhythmically organize isochronous series of external stimuli is found in all humans. It requires that a
subject extracts periodicities from isochronous stimuli and makes periodic temporal anticipation of future events.
Concomitant with these operation, the sensory processing and the speed of perception of stimuli concurrent with the
regular temporal grid is facilitated. It is unknown whether nonhuman primates are also endowed with this perceptual
ability.

Methods
We constructed a tone sequence that was composed of repeating triplets of two short (50 ms) tones and one long (200
ms) tone and that was presented at a constant pulse of 150 tones per minute. This sequence was contrasted with an
irregular sequence of the same number of short and long tones and the same pulse. Human listeners perceptually
organize the regular sequence into triplets with two weak beats followed by a strong beat.

Results
In a first study on humans we show that the different perceptual organizations of regular and irregular sequences are
reflected in different abilities of subjects to detect tones of intermediate duration (100 ms) that are occasionally presented
in place of a long tone. We then show that monkeys that are passively exposed to the same sequences more frequently
changed their facial expressions or their gaze during deviant presentation when the deviants were embedded in a regular
sequence. We also show that approximately one quarter of 175 multiunits recorded from primary auditory cortex of
monkeys fired more spikes, in a particular time range, to regularly presented tones than to irregularly presented tones.

Conclusion
These results suggests that monkeys are capable to hold external stimuli in memory over several seconds. Monkeys are
able to extract periodicities from isochronous stimuli and to make periodic temporal anticipation of future events. Thus the
cognitive ability to hierarchically organize a stimulus sequence appears to be part of the 'natural' and innate behavior of
monkeys. It also implies that some of the characteristics of the music and language faculty in humans may be based on
sensory mechanisms and sensory-motor transformations already present in the last common ancestor of nonhuman
primates and humans.
Concurrent Speech Segregation Based on the Pitch Strength of Iterated Rippled Noise

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Background
In a sound environment with overlapping talkers, voice pitch provides a strong cue to segregation of one speaker among others. We investigated the role of pitch strength in speech-in-background identification for normal-hearing (NH) and hearing-impaired (HI) listeners. Iterated Rippled Noises (IRNs) were created by delaying a noise, applying a gain, and adding this copied noise to the original noise. IRNs produce a pitch that varies in strength with the number of iterations (successive delayed copies added together) as well as with the gain applied to each copy before addition to the original.

Methods
Speech from the Coordinate Response Measure corpus was processed into a series of 16-ms overlapping time windows from which the frequency spectrum was extracted through Fourier analysis. The amplitude spectrum of each window was combined with the phase spectrum of similarly-windowed IRNs and converted back to the time-domain via an inverse Fourier transform. The resulting time waveform contained the frequency contours of speech but the temporal patterns of the IRNs. Target speech was developed with an IRN delay of 10 ms (100 Hz - a voice pitch common for male talkers). Masker speech was generated using two independently generated IRNs, each with a delay of 4.45 ms (224.5 Hz pitch - typical of female speech). Target and masker IRNs were generated with the same number of iterations (2, 4, or 8) and a range of gain factors. The task of the listener was to identify keywords in sentences spoken by the male talker in the presence of the female talker maskers.

Results
For all IRN conditions, identification performance was considerably worse for hearing-impaired subjects than for normal-hearing subjects. However, for both listener groups, better target speech identification was observed for the greater number of iterations, and, to a lesser extent, as the gain increased. Nearly all listeners showed improvements as the pitch strength increased.

Conclusion
Increases in pitch strength of target speech sounds overlapping in time with other talkers results in a consistent and beneficial effect on speech identification for both NH and HI listeners. However, listeners with hearing impairment have great difficulty overall using pitch of IRN to support speech-in-background identification. This is likely a result of the generally weaker pitches produced by IRNs in the impaired auditory system, which may in turn be related to an inability to perceive temporal fine structure. [Work supported by NIDCD]
R113G), impaired calcium binding (CDH23-D101G), subtle weakening of structural stability (CDH23-S47P), or impaired folding (PCDH15-D157G). Interestingly, the different effects of the deafness mutations, studied with biochemistry, correlate with the severity of the reported inner-ear phenotype.

Conclusion
Our results shed light on the molecular mechanics of hair-cell sensory transduction and may help develop tailored treatments for cadherin-mediated deafness.

Noddy, a New Mouse Model for Hereditary Deafness, Illuminates the Function of Protocadherin-15 in Inner Ear Hair-Cell Tip Link
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Background
Considerable evidence now indicates that tip links, structures thought to gate the mechanotransducer channels of the inner ear hair cells, are formed by a tetramer of two cadherin proteins: protocadherin 15 (PCDH15) and cadherin 23 (CDH23), which have 11 and 27 extracellular cadherin (EC) repeats, respectively. Mutations in either PCDH15 or CDH23 often cause inner ear disorders in mice and humans. Using in vitro approaches, we recently showed that PCDH15 and CDH23 bind tip-to-tip in a "handshake" mode that involves EC1 and EC2 repeats of both proteins. However, a paucity of appropriate animal models has slowed our understanding both of the interaction and of how mutations of residues within the predicted interface compromise the integrity of the tip link and consequence of disabling the handshake in vivo. Here, we present noddy, a new mouse model for hereditary deafness and vestibular dysfunction.

Methods
From a recessive N-ethyl-N-nitrosourea (ENU) mutagenesis screen, we identified mice with hearing impairment and abnormal balance. One such line showed lack of an auditory startle response and head-bobbing, and so was named noddy. Various in vivo and in vitro approaches were used to characterize the phenotype and mutation in the noddy mutant mice.

Results
Here we report on a new ENU-induced mouse mutant, noddy, which harbors a missense mutation (I108N) in PCDH15 EC1—the first reported missense mutation in the mouse Pcdh15. Binding of PCDH15-CDH23 fragments is impaired by this mutation in vitro, as indicated by isothermal titration calorimetry experiments. Thus, noddy mutant mice allowed us to test the functional consequences of blocking the handshake interaction in vivo: Noddy homozygotes are completely deaf, mechanotransduction is absent, and hair bundles are fragmented and misoriented. Yet the PCDH15 protein appears to localize normally to the tips of stereocilia. These studies confirm the critical importance of a residue within the putative handshake interface, illustrate, in vivo, how a subtle missense mutation in the tip link can produce deafness and balance disorders, and introduce the first mouse model for this kind of tip-link missense mutation.

Conclusion
The data presented here offer new insights into the interaction between PCDH15 and CDH23, and help explain the etiology of human deafness linked to mutations in the tip-link interface.
TMHS is an Integral Component of the Mechanotransduction Machinery of Cochlear Hair Cells
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Background
Hair cells are mechanosensors for the perception of sound, acceleration and fluid motion. Mechanotransduction channels in hair cells are gated by tip links, which connect the stereocilia of a hair cell in the direction of their mechanical sensitivity. The molecular constituents of the mechanotransduction channels of hair cells are not known.

Methods
To identify components of the mechanotransduction machinery of hair cells, including ion channel subunits, we have analyzed mechanotransduction currents in mouse lines afflicted with hearing impairment. We have also studied the function of the affected proteins by biochemical, cell biological and physiological means.

Results
Here we show that mechanotransduction is impaired in mice lacking the tetraspan TMHS. TMHS binds to the tip-link component PCDH15 and regulates tip-link assembly, a process that is disrupted by deafness-causing Tmhs mutations. TMHS also regulates transducer channel conductance and is required for fast channel adaptation.

Conclusion
TMHS therefore resembles other ion channel regulatory subunits such as the TARPs of AMPA receptors that facilitate channel transport and regulate the properties of pore-forming channel subunits. We conclude that TMHS is an integral component of the hair cells mechanotransduction machinery that functionally couples PCDH15 to the transduction channel.
Activated Radixin is the Major Actin-to-Membrane Connector of Stereocilia
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Background
Radixin (RDX) is a member of the ezrin-radixin-moesin (ERM) family, which serves to organize cell-cortex membrane domains by coupling membrane proteins to actin filaments. Hair-cell stereocilia apparently require at least one ERM member for development. RDX can be activated by sequential binding to phosphatidylinositol 4,5-bisphosphate (PIP2) and phosphorylation; activation unfolds RDX, allowing the C-terminus to bind to actin and the N-terminus to bind to membrane proteins or adapters like SLC9A3R2 (NHERF2).

Methods
We used mass spectrometry to identify and quantify proteins of hair bundles purified from P0 chicken utricles and P21-P25 mouse utricles. We identified and quantified stereocilia structures using electron tomography. We transfected mouse hair cells with in utero electroporation at E12.5, and localized proteins in chicken, mouse, and frog hair cells with immunocytochemistry and confocal microscopy. Protein abundance was assessed with immunoblotting.

Results
RDX was abundant in vestibular stereocilia, present at ~7,000 molecules per stereocilium in chicken and ~4,000 molecules in mouse. By electron tomography, chicken stereocilia had ~7,000 actin-to-membrane connectors. SLC9A3R2 was present at ~30% that level in each species’ stereocilia. In frog hair cells, RDX was found at the periphery of the stereocilia core; in the longitudinal axis, RDX decreased from tapers to tips. An antibody that recognizes activated ERM proteins only recognized RDX above the tapers. The lower edge of the reactivity corresponded exactly to the boundary between the glycosphingolipid (taper) and PIP2 (shaft) membrane domains. In chicken utricle stereocilia, SLC9A3R2 was located in a pattern that was overlapping but not identical to that of RDX. SLC9A3R2 co-immunoprecipitated with RDX extracted from partially-purified mouse stereocilia; binding selectivity of the SLC9A3R2 PDZ domains suggests that it could bind to GPR98 and USH2A, as well as to CDH23 and PCDH15. In mouse cochlear hair cells at P6.5, GFP-RDX was absent from tapers but present in stereocilia shafts, which was confirmed using immunocytochemistry with an anti-RDX antibody.

Conclusion
That localization of GFP-RDX was identical to that of wild-type RDX indicates that we will be able to use mutant RDX constructs to further dissect the role of this protein in stereocilia function. RDX is the only protein present in stereocilia that is present at a high enough concentration, and is in the right place, to be the principal actin-to-membrane connector of stereocilia.

Developmental changes in the mechanotransducer channels of cochlear hair cells and their regulation by transmembrane channel-like proteins
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Background
The first step in sound detection is vibration of the stereociliary bundles of cochlear hair cells to activate calcium selective mechanotransducer (MT) channels. Although the molecular identity of the MT channels is still uncertain, recent evidence indicates that Tmc1 and Tmc2, isoforms of the transmembrane channel like family, play a key role in transduction (Kawashima et al 2011). We have followed these findings by investigating the effects of Tmc knockouts on MT channel properties.

Methods
We examined variations in the hair cell mechanotransducer (MT) channels along the cochlea’s tonotopic axis in wild-type, Tmc1-/− and Tmc2-/− mice between post natal day (P)0 and P10. Patch clamp recordings were made from inner hair cells (IHC) and outer hair cells (OHC) in apical, middle and basal regions of isolated organs of Corti and hair bundles were stimulated with a fluid jet to determine the maximum current. The relative calcium permeability of the MT channel, PCa, was inferred from measurements of reversal potentials in 100 mM external calcium.

Results
MT amplitudes OHCs increased about 1.5-fold from cochlear apex to base but there was little change in IHCs. Similar values were obtained in wild type and mutant mice in the first post-natal week, but after P7, the OHC MT current in the
Tmc1-/- (dn) declined to zero, which may account for the deafness phenotype (Steel & Bock 1980). The calcium permeability of the MT channel, PCa, in different hair cells showed a small but significant decrease from apex to base in OHCs but not in IHCs of wild-type mice up to P6. In Tmc1-/-, PCa in basal OHCs increased to equal that of apical OHCs whereas in Tmc2-/-, PCa was reduced in both apical and basal OHCs. In older mice after P7, PCa in apical OHCs decreased to equal that at the base, but PCa in IHCs was unaltered. In Tmc2-/-, PCa was reduced in IHCs at all ages.

**Conclusion**

We suggest that differences in calcium permeability reflect different subunit compositions of the MT channel which are determined by expression of Tmc1 and Tmc2, the latter conferring higher PCa in apical OHCs and IHCs. Changes in PCa with maturation are consistent with a later down-regulation of Tmc2 in OHCs but not in IHCs. The significance of the transition at about P7, though not understood, is accompanied by other developmental changes in the hair cells. Supported by RO1DC01362

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**Cochlear adaptation is not driven by calcium entry**

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**Background**

To hear, animals rely on specialized mechanosensory hair cells. All hair cells feature a mechanosensory organelle called a hair bundle that is comprised of actin-filled microvilli, termed stereocilia, arranged in a staircase pattern. Deflection of the hair bundle causes activation of mechanoelectrical transduction (MET) channels. MET currents decrease in the face of a constant hair bundle deflection by a process called adaptation, where additional stimulation recovers lost current. Though significant controversy exists over the mechanisms of adaptation, it is universally found that calcium drives adaptation. This data largely comes from frog saccule, turtle auditory papilla, and chick cochlea, with some supporting data from mammalian vestibular and cochlear cells.

**Methods**

Whole-cell patch clamp recording of rat inner and outer hair cells was performed. Hair bundles were stimulated using stiff probes driven by a macroscalepiezo actuator.

**Results**

While further studying the effects of calcium in mammalian cochlear hair cells, we find that rat IHCs and OHCs show robust adaptation at depolarized potentials, when there is no calcium entry. To confirm this result, we performed experiments using different internal calcium buffering conditions and find robust adaptation both when intracellular calcium levels are high or low. Finally, we lowered extracellular calcium with different internal calcium buffering, and again, we find robust adaptation. Shifts observed in activation curves in the presence of low external calcium are independent of internal buffering therefore appear to be due to an extracellular effect.

**Conclusion**

Overall, contrary to accepted adaptation mechanisms, we find that calcium entry is not required for adaptation in mammalian cochlear hair cells.

This work was supported by RO1 DC003896 to AJR, F32 DC010975 to AWP, DAAD Fellowship to TE and NIDCD Core Grant P30-44992.
Audio-visual Interactions in Modulation Discrimination are Modality Specific
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Background
The McGurk and ventriloquist effects attest to the power of cross modal interactions between seeing and hearing. Studies with more abstract stimuli than lips and speech demonstrate that visual information influences sound position, but we know little about how visual stimuli influence the hearing of temporal features in sound and vice versa. We report a unidirectional cross modal influence on participants' ability to discriminate the modulation depths of sounds and shapes.

Methods
In Experiment 1, participants were sequentially presented with pairs of sinusoidally amplitude-modulated sounds and responded whether the modulation depth of each pair was the same or different. Sounds were 250-Hz pure tones modulated at 2 Hz with depths between 20\% and 58\% and duration 1.5s. The modulation depth of each sound pair could differ by 0\% to 32\% in 8\% steps. Sounds were presented alone and in two audio-visual conditions. In the latter conditions, the sound in one interval was accompanied by a cuboid that was modulated about its long axis (rate 2Hz, depth 70\%) in phase with the sound modulation. The cuboid was presented with either the least or the most modulated sound.

In Experiment 2, new participants discriminated the modulation depth of cuboid pairs with depths between 10\% and 18\%. The modulation depth of each pair differed by 0\% to 8\% in 2\% steps. Stimuli were presented alone, or with one interval accompanied by an amplitude-modulated sound (rate 2Hz, depth 70\%). The sound was paired with either the least or most modulated visual stimulus on each trial.

Results
When participants judged the modulation of the two sounds in Experiment 1, the presence of the amplitude-modulated cuboid significantly impaired their ability to discriminate sound modulation depth, relative to the sound-only condition, when the shape was paired with the least modulated sound. There was a trend towards enhanced discrimination when the shape was paired with the most modulated sound. This finding suggests that the visual stimulus enhanced the perceived modulation of the accompanying sound so that participants found it more or less difficult to discriminate sound pairs depending on the interval in which the shape was presented. The presence of the modulated sound in Experiment 2 had no impact on the discrimination of visual shape modulation.

Conclusion
We conclude that the discrimination of modulated sounds is influenced by shape modulation, but that the discrimination of modulated shapes is not influenced by modulated sounds.
Children with Cochlear Implants Demonstrate Working Memory Deficits in Storage, not Processing

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Background
There is growing consensus that hearing loss and consequent amplification likely interact with cognitive systems. A phenomenon often considered in regards to these potential interactions is working memory, modeled as consisting of one mechanism responsible for storage of information in a short-term buffer and another mechanism responsible for processing of that information in some manner. The capability to explore potential interactions between hearing loss/amplification and the operations of this cognitive system has been hindered because there has been no way to index separately the performance of storage and processing. That distinction is important because signal degradation as occurs with a cochlear implant should selectively inhibit the ability to store words in a memory buffer, without affecting processing. The goal of this study was to examine two related hypotheses: (1) Recall accuracy indexes storage while speed of recall indexes processing efficiency; (2) Storage is negatively impacted for children with CIs, but not processing.

Methods
124 children who had just completed second grade participated in this study: 49 with NH, 19 with HAs and 56 with CIs. Listeners heard multiple lists consisting of the same six words (either non-rhyming or rhyming) in different orders, and recalled the order of presentation by tapping pictures on a touch-screen monitor representing each word. Recall accuracy and time required to recall all items were measured.

Results
Three results supported the hypotheses: Recall accuracy for the non-rhyming words was poorer for children with CIs than for those in the other two groups, but was similar across groups for the rhyming words. This outcome suggests that children with CIs are hindered in their abilities to recover phonemic structure from speech signals and use it to store words in a memory buffer. Speed of recall was similar for all children, for both kinds of materials. This finding suggests that the processing abilities of children with hearing loss are not affected by their losses. Recall accuracy and speed were not correlated, reflecting independence of these mechanisms.

Conclusion
It seems possible to measure the operations of storage and processing mechanisms in working memory separately, and that only storage is impaired for children with CIs. This finding could be used in empirical studies to enhance our understanding of the direction and nature of interactions between hearing loss/amplification and cognitive operations by examination of each component of working memory separately.
Investigating Phase Effects in On- and Off-frequency Masking by Schroeder-phase Complexes

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Background

Phase effects in forward masking by Schroeder-phase complexes are thought to be mediated by the interaction of the phase curvature of stimulus with that of the basilar-membrane (BM) filter tuned to the signal frequency, and by BM compression. Although the two factors account well for the phase effects in on-frequency masking, they cannot explain phase effects in off-frequency masking, where the BM should respond linearly. To account for off-frequency phase effects, the activation of the medial olivocochlear reflex (MOCR) has been invoked. Here, the role of the MOCR was tested in two experiments. The first experiment utilized the fact that MOCR activation is stronger for binaural than monaural elicitors to test the prediction that diotic maskers should produce higher masked thresholds, and possibly stronger phase effects, than monaural maskers. The second experiment provided direct measurements of MOCR activation using stimulus-frequency otoacoustic emissions (SFOAEs) with Schroeder-phase complexes as elicitors. A third experiment tested the hypothesis that off-frequency masker phase effects are due to residual BM compression by comparing forward masking produced by on- and off-frequency amplitude-modulated (AM) tones with that produced by unmodulated tones.

Methods

Psychophysical masked thresholds were measured for a 10-ms, 6-kHz signal. The maskers were centered on 6 kHz (on-frequency) and on 3 kHz (off-frequency). The effect of efferent activation on SFOAEs was measured using a continuous fixed-level 6-kHz probe and on- and off-frequency Schroeder-phase MOCR elicitors. Changes in SFOAE due to elicitors were calculated from a heterodyned ear-canal pressure measured in the presence and absence of the elicitor.

Results

Phase effects observed with diotic Schroeder-phase maskers were smaller than those for monaural maskers. The effects of efferent activation by Schroeder-phase elicitors on SFOAEs did not show a systematic dependence on the elicitor phase curvature. Forward masked thresholds measured with an off-frequency AM masker were consistently lower than those for an unmodulated pure-tone masker with the same rms amplitude.

Conclusion

Results from psychophysical and SFOAE measurements do not support the hypothesis that efferent activation contributes to phase effects observed in masking by Schroeder-phase complexes. Results from forward masking by AM and unmodulated tonal maskers are not consistent with phase effects being due to residual compression. Thus, the off-frequency effects of Schroeder-phase maskers remains a phenomenon with no clear explanation.

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Characteristics of Learning an Invented Auditory Non-Linguistic Rule

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Background

Learning of a language involves both implicit and explicit processes that are considered memory-based learning mechanisms. Implicit learning assumes automatic identification of patterns of regularities among similar experiences. Although it does not require conscious effort to discover the underlying rules or structure of the task, it necessitates a critical amount of time and repetition. In contrast, explicit learning involves intentional learning and conscious use of knowledge about events and facts that can occur even after a single exposure. One interesting question is whether these processes are specific to language learning or are they part of a common mechanism that underlie learning of rules with nonlinguistic stimuli. Thus, the goal of the present study was to investigate the characteristics of learning an invented rule using non-linguistic auditory stimuli.

Methods

Ten young adults participated in learning an invented ‘Dolphin language’ in 5 sessions. In each session, each subject listened twice to a modeling list of 16 sequences that consisted each of the same four different sounds (pure-tones and narrow bands) but in a different order according to an invented rule. Then listeners were presented with 16 sequences, half from the modeling list and half were incorrect. Listeners were asked to judge the correctness of these sequences. Listeners were also required to complete sequences. To test transfer of learning, listeners were presented with new sequences and were asked to judge their appropriateness. Dependent variables included percent correct, reaction time (RT) and verbal reports.

Results

Results showed that 50% of the listeners were able to judge correctly the sequences but only a third showed transfer of learning. This suggests that for some listeners implicit learning was involved via memory (and not by discovering the rule). The finding that for some subjects improvement in percent correct was associated with a decrease in RT (supporting implicit learning) while for others with an increase in RT (supporting explicit learning) is in keeping with the notion of variable use of learning processes among subjects.

Conclusion

Although the results of the present study are limited to specific learning conditions, these preliminary data support a common mechanism that underlies learning of linguistic and nonlinguistic grammar. Such information has theoretical as well as practical implications for the diagnosis and intervention of individuals who have difficulties acquiring a language, such as, children with specific language impairment and learners of a second language.
The spatial pattern of cochlear amplification
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Background
Sensorineural hearing loss, which stems primarily from the failure of mechanosensory hair cells, is associated with
changes in the traveling waves that transmit acoustic signals along the cochlea. However, the connection between
cochlear mechanics and the amplificatory function of hair cells remains unclear.

Experiments involving isolated hair cells have identified two force-generating mechanisms. The mechanoreceptive hair
bundles of many tetrapods are capable of generating forces that can be entrained by an external stimulus. Another force-
generating mechanism specific to the outer hair cell of mammals is somatic motility or electromotility: changes in
membrane potential rapidly alter the cylindrical cell’s length. This behavior is mediated by voltage-dependent
conformational changes in the membrane protein prestin.

Although a wealth of evidence suggests that hair-cell forces influence movement of the cochlear partition and vice versa,
it has not been possible heretofore to locally separate the contributions of active hair-bundle motility and somatic motility.

Methods
By applying a scanned-beam laser interferometer in the chinchilla’s cochlea, we recorded the two-dimensional profiles of
the traveling waves elicited by pure tones.

We developed an optical technique that locally and significantly perturbs electromotility. Our technique uses 4-
azidosalicylate, the azide group of which forms covalent bonds with prestin upon activation by ultraviolet light.

To guide our experiments, we developed a mathematical technique that computed a spatial map of cochlear-partition
impedance based on measurements of active traveling waves.

Results
We found that a 500 µm-long segment of the cochlear partition that extended roughly one cycle basal from a wave’s peak,
depending on the location of the hole, encompassed most of the expected region of gain.

We then probed two narrow segments that extended roughly 50 µm along the cochlear partition: one region lying a full
cycle basal to the traveling wave’s peak and another situated just an eighth of a cycle before the peak. Inactivation of the
more basal segment elicited a more gradual accumulation of gain; this caused a small decrement in gain that persisted,
but did not increase, up to the wave’s peak.

Conclusion
These results demonstrate that an active process overcomes viscous damping to locally amplify the cochlear traveling
wave, and that this locally accrued gain accumulates spatially up to the wave’s peak. The results further demonstrate that
prestin plays a crucial role in establishing this gain.
deduced from the cycle width along the force axis. Velocity of the bundle’s tip was varied (0.5-15 µm/s), while the amplitude of motion was fixed (±100 nm). In addition, we studied how the waveform and frequency of spontaneous oscillations were affected by increasing the viscosity of the endolymph that bathed the hair bundles.

Results
At low velocities of bundle motion, friction could be negative, a signature of the active process that drives spontaneous hair-bundle oscillations. When moving at velocities high enough to outrun adaptation, however, we found that friction became positive and displayed a maximum within the narrow region of displacements where the bundle manifested gating compliance, the elastic correlate of transduction channels’ gating. Strikingly, friction was significantly reduced in the presence of a channel blocker (gentamicin). From this reduction, we estimated that channel gating contribute frictional forces 3-5 fold larger than those of hydrodynamic origin. The later were measured after the tip links were severed by applying a Ca\(^{2+}\) chelator (BAPTA). In accordance with these measurements, increasing endolymph viscosity \(\eta\) by thirty-fold had only a mild effect on active hair-bundle movements: oscillation frequencies decreased by only 20% and the speed of the fast transitions within a bundle’s rectangular waveform of oscillation varied like \(\eta^{-0.25}\). A physical description of active hair-bundle mechanics that accounts for the finite activation kinetics of the transduction channels (\(\tau \approx 1\) ms in frog) could quantitatively reproduce the data from both experiments.

Conclusion
We conclude that transduction channels provide a major contribution to hair-bundle friction. Channel properties, but not endolymph viscosity, control damping of hair-bundle movements. Channel friction in turn helps setting the sensitivity and the characteristic frequency of the hair-bundle amplifier.
The High-Frequency Response of an Active Outer Hair Cell is Enhanced by Cochlear Inertia

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Background

Our ears employ an active process that endows them with heightened sensitivity and frequency discrimination. This process results from the action of outer hair cells that amplify the sound-induced motion of the cochlear partition. Outer hair cells perform two forms of movement that potentially contribute to cochlear amplification: active hair-bundle motility and somatic motility. It is unclear, however, how these processes function effectively at high frequencies.

Amplification by active hair bundles has been observed only at low frequencies; moreover, the speed of hair-bundle motion is limited by mechanical loading. High-frequency somatic motility by isolated outer hair cells is severely attenuated by low-pass filtering of the receptor potential driving it, whereas the mechanical load imposed by the cochlear environment inhibits low-frequency somatic length changes.

Methods

To examine the effects of the cochlear environment on the response of outer hair cells at high frequencies, we introduce a model of active outer hair cells. The descriptions of hair-bundle motility and somatic mechanics accord with published observations of movement by isolated outer hair cells. We then use this model to predict the response of an active outer hair cell to sinusoidal forcing.

Results

The model confirms that significant high-frequency motility by an isolated active outer hair cell is unlikely owing to damping and low-pass filtering of the receptor potential. Loading an active outer hair cell with additional damping and stiffness corresponding to that of the surrounding cochlear structures reduces its response even further.

When the masses of the basilar and tectorial membranes are taken into account, however, the system displays high-frequency resonance near the cell's characteristic frequency. The magnitude and sharpness of this response are enhanced when the parameters controlling the activity of the hair bundle are chosen such that the system operates near a Hopf bifurcation, the operating point at which the unforced system transitions from quiescence to spontaneous oscillation. This enhancement by the active hair bundle is possible only when the mass of the tectorial membrane is sufficiently great. Low-pass filtering of the somatic response is counteracted by the activity of the hair bundle.

Conclusion

High-frequency amplification by active outer hair cells is possible owing to a combination of inertial components in the cochlea and the activity of the hair bundle. Somatic motility and active hair-bundle motility collude to create an active amplifier that enhances the motion of the cochlear partition.

Identification of the Voltage Sensing Mechanism in Prestin

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Background

Among the SLC26A proteins mammalian prestin uniquely functions as a direct voltage-to-force transducer of cochlear outer hair cells and the underlying molecule for cochlear amplification. The mechanism of the voltage sensitivity of prestin is still unresolved. So far, two mechanisms are proposed: (1) The extrinsic voltage sensor mechanism is elicited by the intracellular anions, chloride and bicarbonate, that facilitate the voltage-sensor charge movement manifested as nonlinear capacitance (NLC). (2) Positively charged residues within the transporter channel provide the chloride binding site(s) and serve as the intrinsic voltage sensor. We have developed a three dimensional (3D) model of the SulpTP domain (Lovas et al., Proceedings of 22nd American Peptide Symposium, 2011, pp 120-121). The major structural features are 8 transmembrane (TM) spanning domains, two helical pin (HP) re-entry loops and the intracellular finger (IFC). A variant of prestin in which the ICF was removed had substantially reduced NLC. Furthermore, specific residues within the internal anion gating structure and adjacent to the intracellular channel entrance participate in the voltage sensor mechanism of prestin are predicted. These sites were examined to test their contribution to voltage sensitivity.

Methods

Using our 3D structure of the SulpTP domain of rat prestin, chloride binding sites were established and then tested for structure/function relationships in gerbil prestin. Mutations were generated using a QuikChange Lightning site-directed
mutagenesis kit and the resulting nucleotide changes were verified by DNA sequencing. cDNA clones, containing normal and mutated modified gerbil prestin sequences with a C-terminal EGFP tag, were expressed in HEK293 cells. After 24-48 hours of transfection, cells with robust, membrane-associated EGFP expression was used to measure the NLC using whole cell patch clamp technique.

Results
Four tyrosine (Tyr) residues were targeted for substitution with alanine (Ala), phenylalanine (Phe), tryptophan (Trp) and arginine (Arg). All substitutes resulted in a diminished NLC with Tyr>Phe=Trp>>Ala>Arg. Replacement by the positively charged Arg generally caused loss of NLC.

Conclusion
Our observations suggest that a series of Tyr residues, through anion-π interactions, function as intrinsic voltage sensors. These sites are associated with the transporter gating structure and adjacent to the intracellular opening of the transporter pore. These data further suggest that anion-π interactions provide a novel non-covalent binding mechanism by which chloride and bicarbonate anion bind to prestin.

The Slc26 Family Proteins in Membranes Are All Tetramers
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Background
Mammalian prestin, the motor protein of the cochlea outer hair cell, is an oligomer that has recently been shown to be a homotetramer when incorporated into membranes. Prestin is also a member of a large family of membrane proteins, the Slc(Solute Carrier)26 family, which are anion transporters. Although prestin does not transport anions, it requires anions for function, and non-mammalian prestins, transport anions. The sequence similarities of the Slc26 proteins suggest the existence of common underlying mechanisms, including oligomerization. We therefore extended the prior study of prestin oligomerization to determine the stoichiometry of other Slc26 family proteins in membranes.

Methods
We determined the stoichiometry of other Slc26 family proteins, including two non-mammalian prestins and three evolutionarily disparate proteins of the Slc26 family, using the bleach step counting method as previously applied to prestin. Plasmids expressing fusions of Slc26 proteins with enhanced Green Fluorescent Protein (eGFP) were expressed in Human Embryonic Kidney cells. Membrane fragments containing fluorescent Slc26-eGFP molecules were prepared by osmotic lysis. Single fluorescent molecules were isolated and imaged under continuous U-V excitation using an electron-multiplying CCD camera. Bleach step counts were obtained using off-line analysis. A known dimer, Orai1, was studied and analyzed in a similar fashion as a control.

Results
All Slc26 proteins studied were found to be tetramers in membranes.

Conclusion
The existence of tetrameric configurations common to the Slc26 proteins suggests a functional requirement for tetramerization and the existence of conserved oligomerization motifs.
FLIM-FRET Measurements of Voltage Dependent Conformational Changes in Prestin
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Background
Cochlear sensory cells, outer hairs cells (OHCs), elongate and contract in response to changes in transmembrane potential. This electromotility is important for auditory frequency selectivity and sensitivity and is believed to be driven by a motor protein, prestin, which is highly expressed within the membrane of OHCs. While the electrical aspects of prestin function can be studied by measuring the nonlinear capacitance (NLC), there is currently no method to study the mechanical aspects of prestin function.

Methods
We have previously utilized acceptor photobleach fluorescence resonance energy transfer (FRET) to study preston-preston interactions in living cells. However, traditional FRET, based upon detection of fluorescence intensity, is inherently subjected to artifacts that arise from variations in excitation intensity, photobleaching, and fluorophore concentration. We have recently implemented fluorescence lifetime imaging microscopy (FLIM) to perform robust FRET experiments in HEK cells cotransfected with preston-TFP and preston-YFP. FLIM-FRET measurements were performed in patch-clamped HEK cells on the stage of an inverted confocal microscope equipped with a multiphoton laser and time correlated single photon counting electronics.

Results
NLC measurements confirmed that labeled prestin proteins were functional and comparable to wild-type prestin. FRET efficiency was calculated by fitting the lifetime decay of prestin-TFP in cells clamped at various holding potentials. High FRET efficiencies were measured at hyperpolarized potentials, and low FRET efficiencies were measured at depolarized potentials.

Conclusion
The results indicated that changes in the transmembrane potential induce a conformational change in prestin that can be detected by FLIM-FRET. Thus, FLIM-FRET can provide a sensitive assay for the mechanical correlate of prestin function.

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MiRNA183 and MiRNA34a Are Involved in the Onset and Progression of Age Related Hearing Loss
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Background
MicroRNAs (miRNAs), a class of short non-coding RNAs that regulate the expression of mRNA targets, are important regulators of cellular senescence and aging. Our microarray analyses of the organ of Corti from C57 and CBA mice showed that approximately 98 miRNAs exhibited differential expression and that downregulated miRNAs significantly outnumbered upregulated miRNAs during aging. Among these differentially expressed miRNAs, miR34a, a known regulator in the pro-apoptotic pathways, was significantly up-regulated while miR183, known to be important for differentiation and proliferation, was drastically down-regulated. The goal of current study was to examine the temporal and spatial distribution of these two miRNAs during onset and progression of age-related hearing loss (ARHL) and to explore their potential regulatory pathways in the organ of Corti.

Methods
C57BL/6J and CBA/J mice at different time points during aging were used for the studies. Auditory brainstem responses (ABR) thresholds and hair cell morphology were examined to monitor the onset and progression of ARHL. Q-PCR and in situ hybridization were used to quantify and localize the expression of these two miRNAs. Apoptosis qPCR arrays were used to explore target genes of the two miRNAs in the organ of Corti.

Results
Q-PCR analyses showed miR183 was downregulated more than two fold while miR34a was upregulated two fold in both strains of mice during degeneration of the organ of Corti. The changes of expression levels of both miRNAs preceded the
decline of ABR thresholds and hair cell loss. In situ hybridization revealed that miR183 was specifically expressed in hair cells, while miR34a expression in the cochlea was more extensive, in both hair cells and supporting cells. Apoptotic qPCR array analyses showed that many apoptotic genes, such as p53 family, caspase family, Bcl2 family and death domain receptor family members, were drastically altered during degeneration of the organ of Corti in both strains.

Conclusion
miR-183 and miR-34a are differentially expressed during the course of ARHL in both mouse models. Significant up/down-regulation of several major apoptotic gene families suggests these genes might be direct or indirect targets for miR-34a and miR-183. Our study suggests that changes in miRNA expression patterns precede morphological and functional changes and that up-regulation of pro-apoptotic miRNAs and down-regulation of miRNAs important for differentiation and proliferation are both involved in the age-related degeneration of the organ of Corti.

Age effects in discrimination of temporal cues within and between interval groupings of accented tone sequences
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Background
Older persons exhibit difficulty understanding speech that is spoken with an accent and altered timing patterns that differ from their own native language. One change in speech timing results from an elongation of one or more inter-word pause intervals, an outcome that both enhances the perception of accent and creates groupings of longer and shorter sequence intervals. These timing changes could be problematic for older listeners, as age-related deficits in temporal processing are commonly reported. We examine potential age effects on temporal discrimination by using non-speech stimulus sequences that feature some timing attributes of accented speech.

Methods
Reference stimulus sequences consisted of six 50-ms 1000-Hz tone bursts, each separated equally by silent intervals to produce tonal inter-onset intervals (IOI) of 200ms ("Short" or S). Other sequences featured an accent, produced by a lengthening of one or more IOIs to 400ms ("Long" or L). For each of 12 sequence patterns, adaptive forced-choice discrimination procedures were used to measure a duration DL for increments of one targeted sequence IOI (S or L). Listeners included a group of younger normal-hearing adults and two groups of older adults with and without high-frequency hearing loss (N = 15/group). Stimuli were presented monaurally via earphone at 85 dB SPL.

Results
The duration DLs of the two older listener groups were equivalent, indicating that hearing loss did not influence discrimination performance. Younger listeners exhibited better interval discrimination than older listeners for each stimulus sequence. The magnitude of the age-related discrimination deficits was smaller for sequences with the target IOI embedded within similar interval groups (e.g. all Ls or Ss), and substantially larger for a target located between interval groups (e.g., L target between S interval groups).

Conclusion
Comparative discrimination data collected for target intervals within unaccented stimulus sequences indicate that the detrimental influence of sequential accent is much greater for older listeners compared to younger listeners. Diminished temporal sensitivity is evident for target intervals at and adjacent to accent locations within stimulus sequences. Outcomes from these measurements with non-speech stimulus sequences can serve our understanding of the problems experienced by many older listeners when processing the altered temporal cues found in accented speech.
BERA in Aging Grey Mouse Lemurs (*Microcebus murinus*)

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Background
Mouse lemurs are small, gerbil-sized, nocturnal primates, communicating in the high-frequency and ultrasonic range. They are discussed as an emerging new primate model for aging research. The aim of this study was to gain first empirical information on auditory thresholds and hearing sensitivity in mouse lemurs during aging.

Methods
We applied brainstem evoked response audiometry (BERA) a cost-efficient method traditionally used for screening hearing sensitivity in human babies and animal models in hearing research such as cats. Animals were anesthetized using sevoflurane. To assess the effect of age in grey mouse lemurs, we determined frequency dependent auditory thresholds using tone pips in the range between 500 Hz and 95 kHz of two age groups of mouse lemurs (young animals: 1 to 5 years of age; old animals: 6 years or older). Altogether 19 animals were tested.

Results
Audiograms were established individually based on visually determined auditory thresholds. Findings indicate that mouse lemurs show broadband frequency sensitivity from 800 Hz to 40 kHz. Although they exhibit better hearing in the ultrasonic range than most primates, their best frequency of hearing is about 8 kHz. A significant decreased hearing sensitivity over the complete tested frequency range was revealed in aged animals. Furthermore the eldest tested individual was found to be completely deaf.

Conclusion
Long-term measurements will characterize the progress of that possible hearing loss more exactly. Thus, BERA is a promising, cost- and time-efficient technique to estimate hearing capabilities and deficiencies in small primates, such as mouse lemurs.
Age Related Changes in Inhibition in Mouse Auditory Cortex

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Background
Studies attempting to examine age related cortical changes in vivo are often confounded by the dependence of cortical responses upon altered input from the auditory periphery. To obviate this problem, we have employed a slice preparation in which afferents from the auditory thalamus are preserved and can be stimulated in isolation by an electrode to produce responses in the auditory cortex (AC).

Methods
Hearing was measured by ABR at several frequencies (4, 8, 16, 32 and 45KHz) and with Gaussian noise in both young (mean age =7.1 months , n=11) and old (mean age =23.0 months , n=9) CBA/J mice. Auditory thalamocortical slices were cut and cortical responses generated by stimulation of the thalamocortical afferents. These responses were imaged by flavoprotein autofluorescence which utilizes FAD+ as an indicator of metabolic changes induced by neuronal activation. A series of stimulation amplitudes ranging from 5- 500 µA were used, and a 4 - parameter sigmoidal dose -response model was used to fit activation signal in the AC to both an untreated condition and a condition of 150nm GABAzine, a GABA_A- receptor blocker. Basic parameters of the curve in both conditions were extracted and used to evaluate the sensitivity of AC to disinhibition.

Results
The maximum fluorescence above baseline was found to decline linearly with age and hearing threshold in both the normal and GABAzine conditions. Sensitivity to GABAzine was measured as the ratio of maximum fluorescence in the AC during the GABAzine condition, divided by the maximum in the normal condition. Older animals showed a lower sensitivity to GABAzine as indicated by a smaller proportional increase in maximum in the GABAzine condition, which may indicate an age-related decrease in cortical GABAergic inhibition. The differences in old vs. young animals do not appear to be dependent on differences in slice viability, as assessed via AC FAD/NAD ratio, an indicator of metabolic stress.

Conclusion
These data suggest that auditory cortical inhibition in the thalamocortical system may diminish with age and/or hearing loss. More studies are needed to separate out the effects of age and hearing on cortical inhibition.
**Dendritic Extent in Aged Auditory Thalamus**

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**Background**

Age related hearing loss is highly prevalent, occurring in 33% of those over the age of 70 years. Associated with poor central auditory processing, social withdrawal and depression, presbycusis has a major impact on psychological as well as overall physical health. Presbycusis has a multi-factorial etiology, including peripheral deafferentation leading to mal-adaptive changes to the central auditory pathway. The medial geniculate body (MGB) is an obligate auditory brain center in a unique position to gate and code acoustic information. Examining dendritic extent as a measure of neuronal function and processing capacity may be a way to quantify age-related changes in MGB. Early studies found age-related changes in neuronal dendritic extent in neocortex, while more recent studies have not found reduced dendritic extent in aged rat hippocampus.

**Methods**

Neuroleucida imaging and neuron tracing were performed to compare dendritic extent of neurobiotin-filled young and aged Fischer Brown Norway (FBN) rat MGB neurons.

**Results**

Preliminary result from 12 young and 14 age filled MGB neurons showed only small non-significant changes in dendritic extent. The density of the dendrites and the capacitance of the neurons showed no significant age-related changes between MGB neurons: mean dendritic length (young: 4096±317.5 length units; aged: 3948.9±459.6 length units, p=0.80). Surface area also showed no difference between the two groups (young: 14,732.9±1136.7 units, aged: 15,449.5±1895.1 units, p=0.76). Capacitance showed no difference between the two groups (young: 103.1 ± 33.2pF, n=29; aged: 104.6 ± 28.0pF, n=32; p= 0.38).

**Conclusion**

Initial findings suggest small non-significant age-related changes in dendritic extent and whole cell capacitance. Previous studies have shown inconsistent results regarding age-related changes of synaptic structures, dendritic spines and dendritic extent. Fast Golgi Cox studies are underway in order to confirm or reject the present findings.

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**Morphometric analysis and DTI of the auditory cortex in man – changes with aging**

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**Background**

Presbycusis is accompanied in man with changes in the inner ear as well in the central auditory system. The aim of our work was to find out what are the age-related changes in the auditory cortex. Therefore parameters of hearing function were assessed in a group of healthy young controls (YC), a group of elderly with normal presbyacusis (EC) and a group of elderly with expressed presbyacusis (EP) and then their auditory system was examined using Siemens Trio 3T magnetic resonance.

**Methods**

Diffusion tensor imaging (DTI) was performed using EPI sequence. For morphometric analysis and for determination of seed ROI for tractography, 3D structural image was acquired using MPRAGE sequence. Morphometry of the auditory cortex (Heschl’s gyrus – HG and planum temporale – PT) and of the primary visual cortex (V1) was done by means of Freesurfer image analysis suite. Gray matter volume (GrayVol), area of gyral surface (SurfArea) and average thickness of gray matter (ThickAvg) were computed. Probabilistic tractography of auditory pathway from the inferior colliculus (IC) to HG and subsequent DTI analysis was done in FSL environment. Masks of white matter under HG were generated by Freesurfer; masks of IC at both sides of brainstem were defined manually.

**Results**

Significant decrease of ThickAvg in groups EC, EP with respect to YC was found for HG and PT. The analysis showed significantly higher SurfArea on the left side with respect to the right side in HG and PT in all groups. The decrease of the GrayVol for groups EC, EP with respect to YC and higher GrayVol on the left side was in accordance with the ThickAvg
and SurfArea results. Visual cortex showed only a non-significant trend in a decrease of the thickness in both EC and EP. The results from DTI indicated a tendency for increasing L1 in EC and EP with respect to YC in auditory pathway, observable to a greater extent on the right side. No differences between EC and EP were observed in any of the morphological or diffusion parameters.

**Conclusion**

The results demonstrate typical left-right asymmetry of the primary auditory cortex and planum temporale and indicate the degree of the age-related atrophy of these structures that does not depend on the level of hearing dysfunction and is not present to such extent in the primary visual cortex.

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**Age-related changes in sox10 expression in the cochlear lateral wall**

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**Background**

Age-related auditory function deficits may result from a loss of non-sensory cells in the cochlear lateral wall and a decline in the endocochlear potential (EP). Despite the importance of these non-sensory cells, the mechanisms underlying their degeneration remain poorly understood. Sox10 transcription factor is required for the regulation of neural crest-derived cells including several types of non-sensory cells in the inner ear. Mutation of the Sox10 gene causes varying combinations of hearing loss and pigmentation defects in humans. Here, we investigated age-related changes of Sox10 expression in the cochlear tissues of CBA/CaJ mice and human temporal bones.

**Methods**

Young adult (1-3 month old) and aged (2-2.5 year old) CBA/CaJ mice were used because these mice show a greater EP decline at 12 months of age compared to most other mouse strains (Ohlemiller et al., 2010). Sox10 expression patterns were examined in the lateral wall of young adult and aged cochlear tissues obtained from CBA/CaJ mice and human temporal bones using quantitative immunohistochemistry. Ultrastructural changes with age in the cochlear lateral wall also were examined in mouse inner ears using transmission electron microscopy.

**Results**

Nuclear expression of Sox10 protein was present in strial marginal and intermediate cells in both young adult and aged CBA/CaJ mice. Outer sulcus cells and root cells of the spiral ligament also stained positively for Sox10, whereas none of the five types of fibrocytes in spiral ligament reacted positively. Quantitative analysis of Sox10 immunostaining indicated a significant decrease of Sox10+ cells in both the stria vascularis and spiral ligament of aged mice compared to young adult controls. Ultrastructural observations also indicated several signs of lateral wall degeneration in aged mice, which involved multiple cell types in the stria vascularis and spiral ligament. Sox10 immunostaining on the cochlear tissues of human temporal bones are ongoing.

**Conclusion**

Age-dependent morphological and cytochemical changes were seen in multiple cell types in the lateral wall of aged CBA/CaJ mice, an aging animal model with a pronounced EP decline. An age-dependent change in Sox10 gene regulation in the non-sensory cells may contribute the functional decline of cochlear lateral wall and hearing loss in aged mice and humans. This study was supported by NIH R01 DC7506 (H.L.), NIH P50 DC00422 (HL and JRD), NCRR UL1RR029882 and a research grant from the South Carolina Clinical and Translational Research Institute (JRD).

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**Age-associated Changes of Na, K-ATPase Subunit Isoform Distribution and Expression in the CBA/CaJ Mouse Cochlea**

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**Background**

Presbycusis - aging related hearing loss is the number one communication and neurodegenerative disorder of our aged population, and has multi-faceted etiologies. One prominent cause involves declines in the cochlear endolymphatic potential (EP), the cochlear “battery” which is linked to dysfunction of the stria vascularis, The high potassium concentration in the endolymph is maintained by sodium-potassium pumps, such as Na, K-ATPase. It has been reported
that Na, K-ATPase isoforms (subunits) including α1, β1 and β2 are present in the cochlea of young adults, but it is not clear whether their protein expression change with age.

**Methods**

Cochlea were dissected from CBA/CaJ mice: young adults (2-4 months old) and old mice (>30 months). Immunocytochemistry staining with Na, K-ATPase antibodies for α1, β1 and β2 was performed on frozen serial sections of the cochlea.

**Results**

The young adult group displayed strong α1 expression in stria vascularis (SV) marginal cells. The Na, K-ATPase β1 subunit was expressed in SV, spiral ligament, organ of Corti, spiral limbus and spiral ganglion neurons. The Na, K-ATPase β2 staining was observed in SV, organ of Corti and in the spiral ganglion. Comparison of these Na, K-ATPase subunit isoform’s for the two age groups revealed down-regulation of protein expression of all subunits in old mice. It was also noted that atrophies in the aged group, which included a 30-50% shrinkage in cytoarchitectonic area when compared to the young adult group for SV, was confined to the basal part of SV.

**Conclusion**

The main finding of this study is that Na, K-ATPase subunit declines with age and likely plays an important role in age-related hearing loss. Further investigation is needed to clarify the temporal sequence of events for the anatomical atrophy of SV, discharge of the cochlear "battery" and the down-regulation of Na, K-ATPase isoforms. Prevention of these age-related cochlear degenerative events could halt the progression of peripheral elements of presbycusis, leading to novel biomedical interventions to treat age-related hearing loss.

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**Age Related Na+/K+-ATPase Isoform Gene and Protein Expression Changes in the Cochlear Stria Vascularis**

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**Background**

Na+ /K+-ATPase in cochlear marginal cells plays an important role in the stria vascularis’ (SV) function for generation of the endocochlear potential (EP) and maintenance of the ionic composition of endolymph. Isoforms of this ion channel have been reported to localize inside SV including Na+/K+-ATPase α1, β1 and β2 subunits. The effects of aging on these isoforms in SV are not clear. Previously we found that post-translational modification of NKCC1 by Aldosterone may contribute to age-related hearing loss (Ding et al. ARO Abstr. 2012). Since the similarities and functional cooperation between these two ion transport mediators are responsible for generating and maintaining K+ levels in cochlear endolymph, we hypothesize that aging may change Na+/K+-ATPase expression in SV.

**Methods**

Cochleae from CBA/CaJ mice and the cell line SV-K1 (mouse cochlear marginal cells) were used for RT-PCR and Western Blot assays.

**Results**

New RT-PCR experiments revealed that the Na+/K+-ATPase β2 subunit mRNA levels are hardly detected in the cochlea from young adult or aged mice. But relative to young adult mice, Na+/K+-ATPase α1 and β1 subunits’ mRNA levels are down-regulated with age. We then investigated the Na+/K+-ATPase mRNA levels in the marginal cell line. The results were similar to our in vivo studies of mRNA expression levels from cochleae of young adult mice. We then extended our investigation to Na+/K+-ATPase protein expression in vivo and in the cultured cells. Na+/K+-ATPase protein expression is down-regulated in the cochlea of aged mice (30 months old) compared to young mice (2-3 months old). Interestingly, ouabain-treated (an antagonist of Na+/K+-ATPase) SV-K1 cells showed no Na+/K+-ATPase expression changes, suggesting that down regulated Na+/K+-ATPase expression in the aged mouse cochlea may be associated with some so far unknown mechanisms or pathways.

**Conclusion**

To better understand the role of aging on Na+/K+-ATPase down regulation in vivo, further studies are required using Na+/K+-ATPase transgenic mice, along with in vivo measurement of the EP endolymph potential, and measurements of Na+/K+-ATPase promoter regulation assay (Dual-Luciferase® Reporter Assay System) would be helpful to determine if...
increases in Na+/K+-ATPase expression might lead to ion channel activation, moving towards novel biomedical therapies targeted to prevent age-related hearing loss.

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ABR Gap Responses Start to Decline in Middle Age Mice
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Background
The CBA/CaJ mouse strain’s auditory function is normal during the early phases of life and gradually declines over its lifespan, much like human age related hearing loss (ARHL), but on a mouse life cycle “time frame”. This pattern of ARHL is relatively similar to most humans, and currently is not treatable medically. For this reason, CBA mice were used for the present study to analyze the beginning stages and functional onset markers of ARHL. The results from Auditory Brainstem Response (ABR) and Gap-in-Noise (GIN) ABR tests were compared between two groups of mice of different ages, young and middle aged as well.

Methods
For this study, CBA/CaJ mice were used; 10 female and 20 male, and classified into two groups: young adult (Y, N=18, 3 to 4 months old) or middle aged (MA, N=12, 15 to 18 months old). After being anesthetized, ABR GIN ABRs were measured at 80 dB SPL, using wide band noise bursts with gap durations ranging from 0.1 ms to 48.1 msec. ABR tests included an ABR audiogram on each mouse.

Results
For long gap durations the response amplitudes for peak 1 (P1) and peak 4 (P4) to the initial noise burst (NB1) were significantly larger for the Y group. Similarly, P1 and P4 amplitudes to the second noise burst (NB2) were also larger for Y as compared to MA group. P1 amplitudes to NB1 were higher when compared to the NB2 response for the MA group. For NB2, there was a systematic decrease in P1 latency with increases in gap duration for both age groups; however, the NB2 P1 latency was longer for the MA group, particularly for the shorter gap intervals.

Conclusion
At 15 to 18 months old, mice start to show temporal deficits in the GIN test as observed by prolongation of the NB2 P1 latency for shorter gap durations. Effects of ARHL for the MA group were also seen in that the P1 amplitude did not fully recover compared to Y. These findings indicate that age-linked degeneration of the auditory is just beginning in middle age, allowing for biomedical preventative measures to still be a possibility for attenuating further damage due to ARHL.

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Modulation of Mcl-1 expression reduces age-related cochlear degeneration

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Background
Sensory cell degeneration in the cochlea is a major pathology contributing to age-related hearing loss in the adult population. Previous investigations have implicated the apoptotic pathway in age-related sensory cell death. Mcl-1 is an anti-apoptotic member of Bcl-2 family genes that modulates apoptosis-related signaling pathways and promotes cell survival. We have documented the reduction in Mcl-1 expression in aging rat cochleae. Importantly, this reduction in Mcl-1 expression is spatially correlated to the cells undergoing apoptosis. The current study was designed to investigate whether restoration of Mcl-1 expression could reduce aging-related sensory cell degeneration.

Methods
Fischer rats (male and female, 24-25 months) and Sprague Dawley rats (male and female, 2-3 months) were used. Human Mcl-1/enhanced green fluorescent protein plasmid was locally applied to the round window of the rat cochlea. The effect of Mcl-1 transfection on Mcl-1 expression in the cochleae was examined using qRT-PCR, Western blotting, and immunolabeling assays. The auditory function of the animals was assessed using the measurement of auditory brainstem responses. Sensory cell apoptotic activity was assessed with the morphological analysis of the nuclei. The release of cytochrome c from mitochondria, a mitochondrial event of apoptosis, was evaluated using immunohistology.

Results
Strong Mcl-1/enhanced green fluorescence was observed in both sensory and supporting cell regions 15 days after the treatment, suggesting that the in vivo treatment of Human Mcl-1/enhanced green fluorescent protein plasmid transfected both sensory and supporting cells of the cochlear sensory epithelium. Mcl-1 transfection increased both the transactional and protein expression levels of Mcl-1 in aging cochleae. The upregulation of Mcl-1 expression reduced the progression of age-related cochlear dysfunction and sensory cell death. We further demonstrated that the protective effect of Mcl-1 transfection was mediated by the suppression of cochlear apoptotic activity and that the reduction in apoptosis was associated with the suppression of the mitochondrial event of the apoptosis.

Conclusion
The genetic modulation of Mcl-1 expression reduces the progression of age-related cochlear degeneration. Mcl-1 may serve as a potential target for future therapeutic intervention of age-related sensory cell degeneration in the cochlea. (This study was supported by National Natural Science Foundation of China No.30973304 and in part by NIDCD 1R01DC010154)
Age-related Hearing Loss: Gamma-amino butyric acid, Nicotinic acetylcholine and N-methyl-D-aspartate Receptor Expression Changes in Spiral Ganglion Neurons of the Mouse Cochlea

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**Background**
Age-related hearing loss – presbycusis – is the number one communication and neurodegenerative disorder of our aged population. Although speech understanding in background noise is quite difficult for those with presbycusis, there are currently no biomedical treatments to prevent or reverse this condition. A better understanding of the cochlear mechanisms underlying peripheral presbycusis will help lead to future treatments. Objectives of the present study were to investigate gamma-amino butyric acid A (GABA-A) receptor subunit α1, nicotinic acetylcholine (nACh) receptor subunit β2, and N-methyl-D-aspartate (NMDA) receptor subunit NR1 protein expression changes in spiral ganglion neurons of the mouse cochlea, that occur in age-related hearing loss, utilizing quantitative immunohistochemistry techniques.

**Methods**
Frozen serial sections were prepared from young adult (n=6, 2-6 months, 3 males, 3 females) and old (n=6, 28-33 months, 3 males, 3 females) CBA/CaJ mice. The distribution of GABA-ARα1, nAChRβ2 and NMDARNR1 were assayed immunohistochemically. The labeling was measured quantitatively with densitometry using Image J. ABR audiograms confirmed the degree of age-related hearing loss.

**Results**
GABA-ARα1 protein expression decreased with age in spiral ganglion neurons (P<0.05). Surprisingly, the expression of NMDARNR1 increased in spiral ganglion neurons of the aged mice compared with the young adult mice (P<0.01). The nAChRB2 study is ongoing.

**Conclusion**
These findings demonstrated that there are age-related changes of GABA-A and NMDA receptor expression in CBA mouse spiral ganglion neurons, which may be related to functional changes in cochlear synaptic transmission as a function of age. This study leads to more insights into the function of GABA-Aα1, nAChB2 and NMDARN1 receptor protein regulation in the aging mammalian auditory system, paving the way for novel biomedical interventions to prevent or slow down the progression of age-related hearing loss, as well as other types of hearing impairment.

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Noise Exposure and Male–Female Hearing Impairment Differences in Years Lived with Disability in Norway
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Background
Many studies show hearing impairment (HI) as a major contributor to years lived with disability (YLD), yet few are focused specifically on the relative contribution by HI etiology. The Nord-Trøndelag Hearing Loss Study (NTHLS), 1996–1998, affords the opportunity to analyze HI-YLD by sex and etiology based on screening criteria. A population-based sample of 51,574 adults (20–101 years) completed audiometric exams and questionnaires on noise exposure, tinnitus, and hearing health. Engdahl et al. 2005 published audiometric findings contrasting the unscreened and audiologically-screened population, after exclusions based on screening criteria.

Methods
YLD distributions are calculated using World Health Organization (WHO) recommended procedures by 10-year age groups and HI severity: mild, moderate, moderately-severe, severe, profound, and complete. Severity weights are based on AMA-AAO recommendations (JAMA 1979). Subjects are classified by the questionnaire-based screening criteria, including noise exposure history, ear-related disorders, conductive losses, symptoms (tinnitus), and known hearing loss. Noise exposure history included loud occupational exposure, daily exposure from known sources, and frequent exposure to impulse noise. Audiometric criteria associated with noise exposure (Dobie 2005; Hoffman et al. 2006) were used to identify subjects with bilateral audiometric bulges or notches. These two methods were used as proxy measures for the harmful impact of noise exposure.

Results
Age-specific YLD distributions demonstrate that Norwegian men have a steep increase beginning about age 50 and continuing high levels for each subsequent decade of life. Women have a 21% reduction in total YLD due to a delay in the steep increase until age 60. After screening out subjects with noise exposure history, only 49% remain in the sample (males, 25%; females, 75%), which no longer has a sex disparity in HI-YLD. The other screening criteria do not affect the male-female disparity in HI-YLD. If only subjects with bilateral bulges/notches are screened out, then 80% of the sample remains (males, 69%; females, 89%). Screening out these subjects reduces by 60% the excess YLD of males compared to females. Both noise exposure history and bilateral notch/bulge criteria affect primarily HI-YLD of males, with negligible effect on females.

Conclusion
Noise exposure is the major contributor to male-female differences in HI-YLD. For HI and other age-related conditions, delay in age at onset is a preventive measure. Reductions in noise exposure not only reduce total YLD but also diminish age-related sex disparity.
Age-related Changes in Neural Population Representation of Amplitude-modulation in the Presence of Overlapping Maskers

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Background
Age related auditory processing deficits in the central auditory pathway primarily manifest as temporal processing difficulties. Elderly listeners are especially deficient in processing multiple concurrent sound stimuli, or sound stimuli in the presence of partial or complete masking sounds. This study aims to further understand the age related deficits in the neural processing of these sounds in a rodent model of aging.

Methods
Frequency following responses were obtained from young (3-5 months old) and aged (20-22 months old) Fischer-344 rats using subdermal needle electrodes. The target sound stimuli used were sinusoidally amplitude modulated tones (sAM), in the presence of completely or partially overlapping maskers. The maskers used were either broadband noise or sAM tones with the same carrier frequency but a different modulation frequency. The sound levels of the maskers were also systematically varied, to ascertain the growth of masking with level.

Results
As seen in our previous studies, the young exhibited higher FFR amplitudes for the faster modulation frequencies (128, 256 Hz AM) compared to the aged. However, in the presence of other competing sAM tones or broadband noise, the young animals showed a greater degree of relative masking compared to the aged. The masking increased with the degree of overlap, and was the greatest when the maskers were presented concurrently. The decrease in masking with age increased with AM frequency of the target.

Conclusion
This study exhibits the differences in segregation of concurrent sound stimuli, as well as stimuli overlapping with masking noise, with age. The decrease in masking seen with age could, in part, explain the difficulties in processing sounds in the presence of maskers. The decrease in the degree of masking with increase in AM frequency is in concurrence with our previous studies that indicate a decrease in auditory temporal processing with age selectively for faster modulation frequencies. Various explanations for this decrease in masking are consistent with age-related compensatory increases in neural excitability.

Age-Related Changes in the Capillary Basement Membrane of the Stria Vascularis

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Background
Strial presbycusis is one of the 4 predominant pathologic types of presbycusis. Previous studies have shown failure of the basement membrane (BM) structure of strial capillaries in 6- and 12-month-old C57BL/6 mice. The purpose of this study is to examine the cause of age-related changes in the capillary BM of the stria vascularis.

Methods
C57BL/6 mice were grouped according to age as follows: 3 days, 8 weeks, 12 months, and 15 months. Cationic polyethyleneimine (PEI) was used to examine age-related changes in the anionic sites of the BM of the strial vessels. In 3-day-old, 8-week-old, and 12-month-old mice, the bony labyrinths were removed and immersed in a 0.5% PEI solution. Next, they were immersed in 2% phosphotungstic acid and 2.5% glutaraldehyde and embedded in epoxy resin. Ultrathin sections of the cochlear lateral wall were examined using a transmission electron microscope. In 3-day-old, 8-week-old, and 12- and 15- month-old mice, the bony labyrinths were extirpated and immersed in 10% paraformaldehyde and then embedded in paraffin. Serial mid-modiolar 5-μm-thick sections were incubated with either anti-mouse IgG antibody or anti-mouse heparan sulfate proteoglycan (HSPG) antibody.

Results
In 3-day-old mice, PEI particles were evenly distributed on the capillary BM of the stria vascularis and spiral ligament. The area of distribution of PEI particles in the capillary BM of the stria vascularis in 8-week-old mice was smaller than that in 3-day-old mice. In 12-month-old mice, PEI particles were barely detected on the BM capillaries. A statistically significant difference was observed in the distribution of PEI particles between 3-day-old mice and mice in the other groups. Three-day-old mice barely showed any anti-immunoglobulin G (IgG) staining either in the stria vascularis or in the spiral
ligament. In 8-week-old mice, anti-IgG staining was localized in the marginal cells. Further, IgG deposition was detected in the strial capillary BM in 12- and 15-month-old mice. Marked anti-HSPG staining was observed in the spiral ligament, unlike that in the stria vascularis. In the 12- and 15-month-old mice, anti-HSPG staining markedly reduced in the spiral ligament.

Conclusion
An appearance of IgG deposition in the stria vascularis may reduce the anionic sites on the BM of strial capillaries.

Differences in the Processing of Spoken Sentences Between Young and Elderly Normal-Hearing Adults
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Background
Elderly adults experience greater difficulty understanding speech in challenging listening conditions, compared to young adults. This is partially attributed to changes in the auditory periphery which result in loss of audibility. However, even when audibility factors are taken into account, significant ageing effects are still observed. This suggests that changes beyond those in the auditory periphery are also responsible for the challenges faced by elderly listeners. In this project, we investigate the hypothesis that cognitive changes due to ageing also contribute to differences in using context information of spoken sentences.

Methods
Young (M_age = 19.5 years) and elderly (M_age = 66.5 years) participants listened to sentences where the subject either preceded or followed (and was thus semantically constrained by) the verb. Their task was to decide which of four pictures presented on a computer screen was the target, that is, the subject in the sentence they heard. The remaining three pictures represented nouns that were either phonetically or semantically related to the target, or unrelated to it. A desktop eye-tracker recorded the size of the participants’ pupils and the direction of their gaze. Changes in the size of the pupil are used as a measure of changes in the effortfulness of the task, while the participants’ gazes serve as a window to the online processing of speech. The sentences were presented either in quiet or with background, speech-shaped noise.

Results
The two groups showed no significant differences in their scores in the vocabulary and reading span tests, but the elderly participants were significantly slower in a visual speed-of-processing task. Word recognition scores were very high for both groups, with averages ranging between 90% and 98% correct. Semantically constrained nouns were slightly easier to recognize for both groups, but this difference was not significant. The background noise did not have a significant effect on recognition performance, either. The similarities in recognition performance between the two groups ensure that the differences in the way the two groups scan the visual scene are indicative of differences in the way they process spoken sentences.

Conclusion
Based on the preliminary results of this study, we can conclude that eye gazes in this type of paradigm (typically termed “the visual world” paradigm) can be used to shed light on the differences in processing between young and elderly adults. In this study, differences in eye gazes may be attributable to age-related changes in cognitive performance.

Synaptic Aging in the Mouse Cochlea: New Views and Functional Clues
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Background
Studies of sensorineural hearing loss and cochlear compromise have historically concentrated on thresholds and hair cell damage. Although important metrics, recent work has identified widespread loss of IHC-afferent synapses and progressive cochlear nerve degeneration after “reversible” noise-induced hearing loss, i.e. in the absence of hair cell loss
or permanent threshold shift (Kujawa and Liberman 2009). Here we ask if these degenerative changes also appear, with age, in CBA/CaJ mice that are never purposely exposed to noise.

Methods
DPOAEs and ABRs were recorded (5.6 – 44.07 kHz) from an age-graded series (4 to 144 wk) of male CBA/CaJ mice. Immunostained cochlear whole mounts and osmium-stained, plastic-embedded sections (10 µm) were studied at 3 cochlear locations (5.6, 11.3, 32 kHz) by confocal and conventional light microscopy, respectively, to quantify hair cells, ganglion cells and synaptic structures: i.e. synaptic ribbons (CtBP2), glutamate receptors (GluR2) and cochlear-nerve terminals (Na/K ATPase).

Results
Age-related sensitivity loss grows nonlinearly in CBA/CaJ. Threshold elevation is minimal, and confined primarily to mid to high frequencies, out to ~2 yrs. Thereafter, thresholds rise rapidly at all frequencies, particularly for ABR. DPOAE suprathreshold amplitudes are maintained throughout much of the lifespan; in contrast, ABR amplitude declines begin early and progress steadily, even when threshold sensitivity is little changed. Hair cell numbers are maintained until advanced age: OHCs in all rows decline precipitously beyond 2 yrs, first in the apex; whereas, IHC counts change little, even in very old animals. Spiral ganglion losses are modest and slow. In comparison, synaptic changes are dramatic. Ribbons decline throughout life, and throughout the cochlea, falling ultimately to ~50% of young ears. Virtually all surviving ribbons are paired with a glutamate receptor and terminal; thus, the ribbon appears to be a good proxy for the afferent synaptic complex. Moreover, ribbon counts and ABR wave I amplitudes are highly correlated; thus, ABR amplitude is a good proxy for synaptic counts, as long as OHC loss is minimal.

Conclusion
Thus, we document dramatic synaptic aging in the mouse cochlea. Even at ages where threshold sensitivity and hair cells are well preserved, loss of synapses is robust, presumably altering inputs to the aging brain. Results underscore the importance of understanding peripheral status when studying central contributions to auditory aging. Results also suggest that ABR amplitudes may provide non-invasive clues to cochlear synaptopathy, enhancing the possibilities for translation to human clinical characterization. Research supported by NIDCD: R01 DC008577

Effects of Age on the Perception and Neural Representation of Frequency modulation
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Background
Older adults, even with clinically normal hearing sensitivity, have difficulty understanding speech in the presence of background noise. This difficulty may be partly due to age-related declines in the neural representation of sounds.

Methods
Adults with clinically normal hearing sensitivity (thresholds < 25 dB HL at octave frequencies 0.5 – 8.0 kHz) between the ages of 22 and 69 participated in this study. Perception of frequency was tested by obtaining modulation detection limens (FMDLs) at 500 and 1000 Hz using an adaptive procedure. Stimuli were frequency modulated at a rate of 2 Hz. To examine the neural representation of frequency modulation, frequency-following responses (FFRs), which are dependent on phase-locked neural activity, were elicited by FM tones with carrier frequencies of 500 and 1000 Hz. FFR stimuli for each carrier frequency had modulation depths of 0.4 and 2 %. All stimuli were presented at 80 dB SPL. Analysis of FFRs included stimulus-to-response correlations, signal-to-noise ratios, and autocorrelograms. Speech-in-noise understanding was tested using the QuickSIN.

Results
Behavioral FMDLs declined at both 500 and 1000 Hz as age increased. FFR stimulus-to-response correlations and autocorrelograms showed greater age-related declines at 1000 Hz than at 500 Hz. FMDLs and FFR measures were correlated with speech-in-noise understanding.

Conclusion
These results suggest that the perception and neural representation of frequency modulation is negatively affected by age, even in the absence of significant hearing loss. [Supported by the National Organization for Hearing Research Foundation]
Single Unit Recordings in the Cochlear Nucleus of the Anesthetized Rat
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Background
Neuronal types in the cochlear nuclei can be distinguished on the basis of both physiological and morphological criteria. The neuronal types participate in a complex neural circuitry with both excitatory and inhibitory components and project to higher auditory nuclei. Frequency response areas (FRA) and peri-stimulus time histograms (PSTH) have been used to characterize different neuronal populations in several species, but response types in the rat have not been described. Here, we describe FRA and PSTH patterns recorded in the same neurons in the DCN of the anesthetized rat.

Methods
Experiments were performed on 17 female Long-Evans rats anesthetized with urethane (1.5 g/kg, i.p.). Glass micropipettes filled with 2 M NaCl (15–25 MΩ) or tungsten electrodes (1–2 MΩ) were used to make extracellular recordings of neural responses in well-isolated single units. Electrolytic lesions (10–15 µA for 10–15 s) were made after recording and used for subsequent histological verification of the recording sites. Auditory stimuli consisted of 35 ms long pure tones with a 5 ms rise/fall ramp at a repetition rate of 4 Hz.

Results
The present data set includes the response profiles of 45 single units in the cochlear nuclei. Twenty-seven of these were localized to the DCN and will be described here. (Eleven units were localized to the VCN and 7 could not be assigned a location.) The recorded neurons include a wide variety of firing patterns and rate-level functions, as have been described in detail for other species. More than the half of the neurons in the DCN (17/27) displayed non-monotonic rate-level functions. Based on the PSTHs, DCN neurons were classified as, primary-like (n = 12), pause/build (n = 6) and onset (n = 9) firing patterns. According to FRAs, neurons were identified as type II (n = 16), II/III (n = 8). We found the 3 PSTH types for type III FRA, primary-like and onset firing pattern for type II/III, and primary-like was found to be only type II FRA. In the VCN, our small sample included the primary like, n = 5; pause/build, n = 2; onset, n = 3 and chopper, n=1.

Conclusion
Our study describes basic temporal and spectral properties of unit responses in the DCN of the anesthetized rat.

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Synergistic Action of GABA and Glycine Provides Powerful Inhibition to Spherical Bushy Cells in the Cochlear Nucleus of Gerbil
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Background
In the auditory brainstem nuclei of mature rodents, precisely timed inhibition is predominantly mediated by the fast glycnergic neurotransmission which contributes to accurate timing of action potentials. GABA is transiently released from inhibitory terminals during early postnatal development (LSO), but in the mature system it mainly plays a modulatory role through presynaptic GABA_B receptors (MSO, AVCN) or postsynaptic glycine receptors (MNTB). In the AVCN, presynaptic terminals containing glycine and/or GABA and respective postsynaptic receptors are present not only during development, but also in maturity. Although in vivo recordings with iontophoretic drug applications consistently suggested a contribution of both transmitters to the inhibition on spherical bushy cells (SBCs), our understanding of their action and the advantage of such a two-transmitter based system beyond development are still inconclusive.

Methods
Physiological effects of inhibition were assessed by means of in vivo extracellular recordings of SBCs, conducted in P20-30 Mongolian gerbils using either a sole inhibitory stimulation (stimulus within the unit’s inhibitory sideband) or a two-tone stimulation protocol (additional excitatory stimulus at unit’s CF). In slices containing AVCN, pharmacologically isolated, evoked inhibitory postsynaptic currents (eIPSCs) and potentials (eIPSPs) were recorded in SBCs at postnatal days P22-
30. The contribution of glycine- and GABA-mediated synaptic transmission was measured following application of GABA_A^- and glycine-receptor antagonists. Amplitudes and waveform kinetics of inhibitory currents (glycinergic, GABAergic and synergistic) measured in vitro were used to constrain a conductance based model of synaptic interaction in SBCs.

Results
Acoustically evoked inhibition changes the ratio of isolated EPSP/APs and prolongs the EPSP-AP transition. In slice recordings, eIPSCs exhibit slow synaptic decays causing a stimulus number- and frequency-dependent conductance plateaus. Glycinergic transmission largely determines the amplitude and the kinetic of eIPSCs and eIPSPs, while the contribution of GABA_A^- receptors potentiates the AP inhibition. The amplitude of eIPSCs is larger than the sum of pharmacologically isolated glycinergic and GABAergic components. Physiological significance of synergistic action is suggested by the modeling results, showing the increased inhibitory potency at a wider range of excitatory conductances and longer duration of inhibition.

Conclusion
Efficient synaptic inhibition of SBCs is mediated by a synergistic action of glycine and GABA. The joint action of transmitters causes a supra-additive increase of the dynamic range in which the inhibitory conductance prevents the AP generation by the excitatory synaptic input. Still, the physiological mechanism underlying a possible interaction remains to be elucidated.

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Neural Correlates of Tone Detection in the Cochlear Nucleus of Nonhuman Primates
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Background
Detection of sounds is a critical and fundamental function of the auditory system. Our behavioral studies showed that thresholds to detect a tone displayed a characteristic U-shaped relationship with frequency. Reaction times to detect tones decreased as the tone sound pressure levels increased beyond threshold. To evaluate the neuronal correlates, we measured the responses of neurons in the inferior colliculus (IC) to tones. Signal detection theoretic analysis revealed that neuronal thresholds and slope of neurometric functions in IC matched behavioral thresholds and slope of psychometric functions. The responses of single units were different on correct trials relative to incorrect trials at the same signal sound pressure level. The behavioral reaction times could be related to the magnitude of IC neuronal responses. These results suggest that IC neuronal response strengths were strongly correlated with behavior. To test the hypothesis that the response correlations were derived in the IC and were not inherited from the responses of the cochlear nucleus (CN), we simultaneously measured behavioral responses and neuronal responses in the CN, to evaluate the role of the CN in detection.

Methods
Two macaques (Macaca mulatta) were trained in a reaction time lever release task to report the detection of tones. Appropriate catch trials were interleaved to make sure the monkeys were actually reporting tone detection. Single units were simultaneously recorded from the CN (n=60). The characteristic frequency of these units ranged up to 32.5 kHz. Most of the units were classified as type I or type I/III or type III, based on the distribution of excitation and inhibition in frequency response maps. Signal detection theory and Receiver Operating Characteristic (ROC) analyses were used to create psychometric (behavioral), and neurometric functions. Neurometric functions reveal the probability of correct response based on the distribution of the number of spikes fired by the CN neuron on tone trials and catch trials.

Results
On the average, CN neuronal thresholds were larger than behavioral thresholds by about 19 dB. In addition, the slopes of the neurometric functions were shallower than those of the simultaneously measured psychometric functions. The responses of some neurons were different based on the behavioral outcome of the trial, but the number of these was smaller than in the IC.

Conclusion
These results suggest that the magnitude of CN responses are only weakly correlated with behavior, and a model is suggested to transform CN responses into IC responses. Supported by R01 DC11092.
Modulation of Temporal Coding in the Superior Paraolivary Nucleus
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Background
Precise temporal coding by auditory system neurons is essential to accurate hearing. The superior paraolivary nucleus (SPON) is a GABAergic nucleus of the superior olive whose neurons are particularly sensitive to discontinuities in auditory stimuli. It is known that the characteristic offset spikes of SPON neurons result from a post-inhibitory rebound mediated by strong glycinergic inputs from the medial nucleus of the trapezoid body. However, less is known about the modulatory effects of excitatory inputs (originating from the cochlear nucleus) or GABAergic inputs (arising from collateral innervation from other SPON units) that also impinge on SPON cells.

Methods
SPON neurons were manipulated pharmacologically in vivo in anesthetized adult female rats. “Piggyback” electrodes, in which a recording electrode was glued to a multibarrel glass pipette, were loaded with either the glycine antagonist strychnine, the GABA antagonist Gabazine, the AMPA receptor antagonist NBQX or the NMDA receptor antagonist CPP. Single SPON units were isolated and characterized using a battery of auditory stimuli, including stimuli designed to measure the response of the units to sound gaps and sinusoidally amplitude-modulated tones. Drugs were then administered iontophoretically, either separately or in combination, and the units were monitored to detect differences in firing patterns.

Results
As previously reported, only blockade of glycine receptors greatly diminished the typical offset spiking patterns to pure tones. Moreover, when glycine neurotransmission was blocked SPON neurons showed both increased spike rates and decreased synchronicity to amplitude modulated tones, as well as altered responses to sound gaps. Blockade of glutamate receptors reduced the ability of SPON neurons to synchronize with modulation rates above 200 Hz, whereas blockade of GABAergic inhibition showed little effect on temporal processing. We are currently investigating the functional role of glycine, GABA and excitatory neurotransmitters in modulating other aspects of SPON response properties.

Conclusion
Preliminary data suggests that blockade of glycine, GABA-A or glutamate receptors caused changes in the ability of SPON neurons to phase lock to increasing rates of amplitude modulations. This finding indicates that each of these neurotransmitters has a role in modulating aspects of the temporal response properties of SPON neurons.

Contribution of NMDA and AMPA Receptors to Response Properties of Neurons in the Medial Nucleus of the Trapezoid Body of the Rat
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Background
The medial nucleus of the trapezoid body (MNTB) is a prominent cell group of the superior olivary complex that plays a critical role in sound localization circuitry. MNTB principal cells, the dominant cell type in the nucleus, receive their excitatory input from globular bushy cells of the contralateral cochlear nucleus via large calyces of Held. While the excitatory input to MNTB neurons carried via the calyx of Held is mediated by fast-acting AMPA receptors (Banks and Smith, 1992), NMDA-mediated excitation with a slower time course has also been documented in rats younger than postnatal day 16 (Forsythe and Barnes-Davies, 1993; Hamann et al., 2003) and mice (Joshi and Wang, 2002).

Methods
In present study, single units in the MNTB were manipulated pharmacologically using “piggyback” electrodes in vivo in anesthetized adult rats. We compared contributions of NMDA and AMPA/kainate receptors to auditory evoked responses in MNTB. We examined response magnitude, temporal patterning, gap detection threshold (GDT) and responses to sinusoidally amplitude modulated (SAM) tones before, during and after local micro-iontophoretic application of the NMDA receptor antagonist CPP [(±)3-[2-carboxyipiperazin-4-yl]-propyl-1-phosphonic acid], the AMPA receptor antagonist NBQX [(1,2,3,4-tetrahydro-6-nitro-2,3-dioxo-benzo[ff]quinoxaline-7-sulfonamide], or both drugs applied simultaneously.

Results
CPP and NBQX reduced spontaneous and evoked firing rates, and resulted in longer first-spike latencies and increased jitter in the timing of the first spike. The effects of NBQX and the NBQX/CPP cocktail were much stronger than CPP...
administered alone. For some neurons, GDTs were markedly increased by application of NBQX. Each drug also reduced the firing rate and entrainment of MNTB neurons in response to SAM tones, but did not change vector strengths.

**Conclusion**
Overall, blockade of AMPA receptors was more effective in modulating response properties of MNTB neurons than blockade of NMDA receptors. However, the results obtained under pharmacological blockade of the NMDA receptor suggest that NMDA-mediated excitation at the calyx-MNTB synapse persists into adulthood.

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**The temporal responses of single units in the ventral cochlear nucleus to conspecific vocalisations**

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**Background**
The neural mechanisms responsible for the difficulties in understanding vocalisations in adverse listening conditions are still debated. Here we compare the representation of temporal fine structure (TFS) and envelope (E) cues of complex sounds, including conspecific vocalisations, in the responses of single units from the ventral cochlear nucleus of normal-hearing guinea pigs with the responses from guinea-pigs with a mild sensorineural hearing loss.

**Methods**
Single units have been recorded from the ventral cochlear nucleus (VCN) of 7, normal-hearing, urethane anaesthetised guinea pigs. For a further 7 animals we produced a moderate sensorineural hearing loss by exposing the guinea pigs to a 2 kHz sinusoid at 120 dB for 1 hour. The 2-kHz region was selected to lie within the phase-locking limits of the guinea pig. This exposure resulted in a threshold loss of about 20 dB between 0.1 and 3 kHz and less than 10 dB loss between 3 and 7 kHz. Recordings from animals with a hearing impairment commenced around 30 days post-exposure. The stimuli were pure tones (for comparison with previous studies) and both conspecific and non-conspecific animal vocalisations. We selected signals to provide a variation in temporal envelope and spectral range. Four guinea pig vocalisations were studied; the chirp, purr, chutter and whistle. All vocalisations were randomly presented for a total of 50 repeats.

**Results**
We have recorded from 80 units in animals with normal hearing and 81 units from animals with a mild sensorineural hearing loss. As expected many units in the VCN produced a good temporal response to the waveform of conspecific sounds. This response was correlated with unit type with primary-like units showing the best response to the temporal aspects of the sound although the onset units showed a strong response to the peaks of the temporal envelope. In animals with a mild sensorineural hearing loss the thresholds of fibres in the 1-3 kHz region were raised by approximately 20 dB re: thresholds in our normal hearing population. Although thresholds were elevated we found no significant difference in the tuning between normal and hearing impaired animals.

**Conclusion**
A preliminary analysis of the data suggests there is little change in the temporal discharge properties to conspecific vocalisations following a mild sensorineural threshold loss. This is consistent with the small shift in unit thresholds and corresponding lack of significant changes in tuning.
Stability and Plasticity of Brainstem Function Across the Lifespan
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Background
The general consensus is that the human auditory brainstem undergoes rapid developmental changes early in life (until age ~2) followed by a prolonged period of stability that terminates when aging-related changes in brainstem function emerge. However, earlier work on human brainstem development was limited by inadequate sampling across the lifespan and/or averaging across children and adults. Using a larger dataset than past investigations, we aimed to trace more subtle variations in brainstem function that occur normally from infancy to the 7th decade of life.

Methods
We recorded auditory brainstem responses (ABRs) to a click and the speech syllable "da" in normal-hearing healthy individuals between the ages 0.25 to 70 years. Suprathreshold stimuli were presented monaurally to the right ear. Analyses were performed on a dataset of 580 subjects, divided into 13 sex-matched age groups. We measured the latency of each prominent peak in the speech- and click-ABR. For the speech-evoked ABR, we also measured frequency encoding across the response spectrum, in addition to gauging the trial-by-trial consistency of the response.

Results
Although each family of measures (latency, frequency encoding, response consistency) has a distinct developmental profile, there is a common trend for developmental changes to continue well past age 2. In the case of timing and frequency encoding, the developmental profile is characterized by a relatively transient developmental apex around ages 8-10. That is, latencies become progressively earlier and frequency encoding becomes progressively more robust until the apex, after which latencies become longer and frequency encoding declines until early adulthood where the developmental trajectory stabilizes. In the case of response consistency, the developmental apex is more prolonged, with trial-by-trial consistency increasing from infancy to age 8 and then staying relatively stable into early adulthood. Across all measures, aging-related changes begin to emerge in the fifth decade of life.

Conclusion
This study provides the first comprehensive assay of how the ABR changes over the human lifespan. The outcomes of this research provide new insight into auditory development. It establishes that developmental plasticity continues well into the second decade of life, which challenges conventional wisdom that the auditory brainstem is mature by age ~2. We interpret the developmental apex as reflecting a sensitive period in development when experience, including auditory remediation and deprivation, may be most influential.

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www.brainvolts.northwestern.edu
The mouse anteroventral cochlear nucleus (AVCN): Regional differences in synaptic organization of bushy cells

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Background
Spherical and globular bushy cells (SBCs and GBCs) of the AVCN receive huge auditory nerve endings, the endbulbs of Held and modified endbulbs, which are specialized for precise representation of acoustic information. These high fidelity synapses are essential for locking neural activity to acoustic events to encode features such as interaural level and timing differences used to compute sound location. Recent physiological studies in mice and other species have called into question the clear distinction between SBCs and GBCs, suggesting instead a sort of continuum of response properties determined by the inputs to the cell with exemplars of both response types at either end of the spectrum. We conducted a systematic investigation of mouse bushy cells along the rostral-caudal axis in an effort to more completely understand the morphological variation that gives rise to reported response properties in mice.

Methods
We combined quantitative light and electron microscopy to investigate regional differences in the AVCN of CBA/CaJ mice. Cell morphology, immunostaining, and the distribution of primary and non-primary synaptic inputs along the rostral-caudal axis were characterized. Morphometric analysis was accomplished using ImageJ.

Results
Overall, large regional differences in postsynaptic cell characteristics were not found. In contrast, bushy cells of the rostral AVCN received a greater proportion of axosomatic input than bushy cells of the caudal AVCN. The percentage of terminals displaying primary auditory nerve morphology was larger in caudal AVCN, while non-primary excitatory and inhibitory inputs were more common in rostral bushy cells (Figure 1). The ultrastructural characteristics of primary auditory nerve inputs revealed many similar characteristics in rostral and caudal AVCN (Figure 2). Primary terminals were large, replete with synaptic vesicles, and often displayed multiple postsynaptic densities and structures associated with ultra-fast auditory synapses (mitochondrion adherens complexes, extended extracellular spaces). Cross sectional area, postsynaptic density length and curvature, and percent mitochondrial area were not different in axosomatic auditory nerve terminals in rostral and caudal AVCN; however, rostral auditory nerve terminals contained more synaptic vesicles near the postsynaptic densities. Primary axosomatic auditory nerve terminals also formed synapses with adjoining dendrites.

Conclusion
Regional differences in physiological responses of mouse bushy cells likely depend mainly on the pattern of inputs rather than morphological characteristic of the cells themselves. These findings suggest that rostral bushy cells may display greater postsynaptic synaptic integration than caudal bushy cells, but further characterization is needed.
Microglial Cells Participate in Re-Shaping Neuronal Circuity of the Rat Auditory Brainstem After Cochlear Ablation

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Background
Cochlear ablation triggers a wealth of cellular and molecular responses in the central auditory system of adult rats. Axonal degeneration, neuronal shrinkage, astrocyte hypertrophy, and axonal growth related to the expression of the growth-associated protein GAP-43, lead to complex rearrangements in auditory brainstem circuitry. Microglia are known to phagocyte cellular debris, secret cytokines and growth factors, and remodel the extracellular matrix in neuropathological conditions. Although previous work has shown a reaction of microglia following the intracerebral injection of heat-killed bacteria, visual-experience, axotomy, and cochlear ablation, only little is known about their participation in synaptic plasticity. We here asked whether microglia respond to sensory deafferentation with identifiable actions on central neuronal circuitry.

Methods
Young adult Wistar rats were unilaterally deprived from primary auditory input by cochlear ablation. After survival times of one to seven days, immunocytochemistry for light and electron microscopy was performed. Double-labeling was done for several combinations of cytological markers, signaling molecules, and second messengers. This approach allowed us to study the spatial relationship between microglia identified by CD11b and neuronal compartments before and after sensory deafferentation.

Results
By one to seven days post-lesion, activated microglia have developed intense immunoreactivity for MAP kinases p-ERK1/2 and p-p38 in ventral cochlear nucleus (VCN), lateral superior olive (LSO), and central inferior colliculus (CIC). Labeling for p-ERK1/2 and p-p38 was largely absent outside microglial profiles, with the exception of selected presynaptic endings. Remarkably, these microglia were found in close apposition to neuronal somata, often engulfing presynaptic endings. Most of these endings contained either vGluT1 or GAD65. Microglial processes showed morphological signs for phagocytic actions on synapses, suggesting that they participate actively in altering the architecture of neuronal networks. This was apparent not only in VCN directly affected by axonal degeneration, but also in LSO and CIC where modification of the pattern of synaptic contacts must be invoked by secondary causes. Furthermore, the microglial reaction was seen in AVCN and LSO to precede astroglial hypertrophy, supporting the concept of microglia-derived cytokines activating astrocytes.

Conclusion
In central auditory pathways, microglia displaying MAPK-signaling appear to contribute to an adaptive response to sensory deafferentation. This response occurred in regions that were directly affected by axonal degeneration as well as in others only indirectly affected, and appears to include two ways of action: (1) activation of astrocytes, and (2) synaptic rearrangement of neuronal networks.
Synaptic Organization of Posterior Ventral Cochlear Nucleus (PVCN) Magnocellular Core in the CBA/CaJ Mouse

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Background
Large cells of the PVCN are thought to be essential for many fundamental auditory processing tasks such as loudness perception, listening in noisy backgrounds, spectral and temporal integration, and modulation of the contralateral cochlear nucleus. Much of what we know about the hypothesized roles of PVCN cells comes from studies performed in mice, yet most of what we know about the ultrastructure of PVCN synapses comes from studies performed in cats and rats. A detailed characterization of synaptic organization of PVCN in mice is needed to enhance our understanding of the role of synaptic organization in generating specific physiological responses and to accurately identify age-related, environmental, and genetic contributors to hearing loss that may affect auditory processing in PVCN in mouse models.

Methods
We used electron microscopy, immunohistochemistry, light microscopy, and unbiased stereological methods to reveal qualitative and quantitative details regarding the baseline state of synaptic organization of the PVCN magnocellular core in young adult CBA/CaJ mice.

Results
The CBA/CaJ PVCN magnocellular core contains approximately 110,000 to 120,000 VGLUT1-positive auditory nerve puncta within approximately 110 to 120 million µm³ as measured using stereological probes. Prototypical PVCN neurons are depicted in the figure (scale bars = 2 µm). Globular bushy cells with oblong or round somata resembled those of the AVCN with auditory nerve (blue), inhibitory (red) and non-primary excitatory inputs (light green) contacting most of the soma. T stellate cells received few axosomatic terminals, and much of their somata were ensheathed in glia. Many small auditory nerve boutons, inhibitory terminals, and nonprimary excitatory terminals contacted their dendrites. T stellate cells were often adjacent to other T stellate cells, but membranous specializations suggestive of junctions were not obvious. D stellate cells with variable soma shapes received numerous, mostly inhibitory axosomatic terminals. Small auditory nerve boutons and non-primary excitatory terminals were also observed. D stellate dendrites also received many inputs. Auditory nerve terminals of varying size covered much of the octopus cell somata and dendrites. Specializations common to other fast auditory synapses were observed in some of these terminals. Non-primary excitatory terminals were also observed (dark green), possibly originating from axon collaterals from other octopus cells.

Conclusion
The general synaptic organization of the PVCN magnocellular core is similar to other species. These data will form the basis for future quantitative studies of PVCN neural plasticity and degeneration in response to genetic and environmental manipulations and aging.
Laminar and cellular organization of the human dorsal cochlear nucleus
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Background
Tinnitus, the perception of a phantom sound, or “ringing in the ears” is a debilitating condition affecting as many as 10% of the US population. It is usually, but not always, associated with hearing loss. In some patients, tinnitus may be modulated by somatosensory input (somatic tinnitus), suggesting somatosensory-auditory interactions. Much work has been devoted to determining the CNS site of tinnitus generation. Especially useful have been studies using behavioral animal models of tinnitus, usually rodents. Data from these studies have implicated the dorsal cochlear nucleus (DCN) as a key site in tinnitus generation. In nonprimate mammals, the DCN has a distinct laminar organization and several anatomically and neurochemically defined neuronal types. The circuitry in the nonprimate DCN has been well-characterized. Studies of rodent models of tinnitus have shown an increase in spontaneous activity in the projection neurons (fusiform or pyramidal cells) of the DCN. Further, anatomical data in rodent have shown that the DCN receives both auditory and somatosensory information, suggesting it as a mediator of somatic tinnitus. However, there is a major problem in extending the findings from the rodent DCN to humans. Classic studies of human DCN suggest that its structure is substantially different from that of the rodent, with an erosion of laminar organization, and the loss of several cell types, including granule cells. These results suggest major differences in circuitry in human DCN compared to that in rodents and question the usefulness of data from rodents in understanding tinnitus in humans.

Methods
We have reexamined the organization of the human DCN using cases from the Witzelton Normal Brain Collection. We used Nissl-staining and immunohistochemistry for several markers including nNOS (nitric oxide synthase), calretinin, calbindin, nonphosphorylated neurofilament protein and VGLUT2 and VGLUT1, markers that label axon terminals of somatosensory and auditory afferents in the rodent DCN.

Results
Our data show that the human DCN has three distinct layers, an outer molecular layer, a layer of large projection neurons and a deeper layer with multiple cell types. We also found axon terminals immunoreactive to VGLUT2 and VGLUT1, suggesting both somatosensory and auditory inputs to the human DCN.

Conclusion
Contrary to earlier studies, our results suggest that the organization and inputs of the human DCN are not that different from the rodent DCN and support the usefulness of rodent models in studying the role of the DCN in the generation of tinnitus.
3D reconstructions of dorsal cochlear nucleus neurons and their synapses using Serial Block Face Scanning Electron Microscopy
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Background
Serial Block Face Scanning Electron Microscopy (SBFSEM) provides a rapid, reproducible means of viewing ultrastructural features from biological samples in 3D. Using SBFSEM for reconstructing ultrastructural details of synapses and synaptic circuitry in the dorsal cochlear nucleus (DCN), we sought to reconstruct and map synaptic terminals on different regions of fusiform cells, the principal output neurons of the DCN.

Methods
Fixed and stained samples of the DCN were embedded in epon then sliced and imaged on a Zeiss Sigma VP scanning electron microscope equipped with a Gatan 3View stage/knife system. The sections were cut at 75 nm thickness in the transverse plane at a level where the DCN reaches its maximal dorsoventral dimension. Images were obtained in 6 adjacent fields, which together, spanned the molecular, fusiform cell and outermost part of the deep layers of the DCN. In each field, a stack of over 600 sections was cut and imaged such that the total thickness of each stack was approximately 50 um. After image acquisition, the resulting stacks were stitched together into a montage that measured 160 x 240 um. Within this montage, the outlines of fusiform cells and their dendrites and inputting synapses were traced.

Results
The cells were classified as fusiform cells if they met the following criteria: soma measured approximately 25-30 um in diameter, dendritic arbors extended toward both the ependymal surface (apical dendrite) and deep layer (basal dendrite), an axon extended toward the dorsal acoustic stria and the apical dendritic arbors were studded with spines. To obtain insight into how synapses are spatially distributed over the fusiform cells, we mapped the terminals over the somal surface and on selected segments of their primary and secondary dendrites. We also obtained measures of the number and sizes of dendritic spines on the apical dendrites. In selected subsets of terminals, the number of neurotransmitter vesicles and dimensions of synaptic densities were quantified. Lastly, we attempted to trace axons from their synaptic ending to determine whether axons can be traced to their cells of origin and to assess whether different synaptic terminals arise from different branches of the same axon.

Conclusion
We are using this data to establish normative values that can be used for comparison with similar measures obtained from animals that have been exposed to intense sound. (Supported by NIH R01 DC009097).

Suppression of Neural Firing in the Inferior Colliculus Induced by Focal Electrical Stimulation of Auditory Cortex: Implications for the Treatment of Tinnitus
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Background
One potential treatment option for tinnitus suppression utilizes cortical electrical stimulation to reverse abnormal neural changes causing the tinnitus percept. Although initial clinical trials have been encouraging, cortical stimulation results have varied dramatically across patients. In order to improve clinical results, we sought to determine the neurophysiological effects of focal electrical stimulation of auditory cortex (AC) on the auditory pathway. Particularly, we assessed whether AC stimulation can suppress neural firing within the inferior colliculus (IC), a region that has shown hyperactivity in tinnitus animals and patients. Since primary auditory cortex (A1) is difficult to surgically access in humans (hidden within Heschl's gyrus), we focused the majority of our investigation on secondary auditory cortex (A2).

Methods
We positioned multi-site electrode arrays into AC and the IC of ketamine-anesthetized guinea pigs. After characterizing the acoustic-driven responses on each site to confirm its location (i.e., frequency region and sub-nuclei of AC or IC), we electrically stimulated the deeper output layers of AC and recorded the corresponding changes in spontaneous and acoustic-driven responses throughout the IC.

Results
Stimulation of deeper A1 output layers suppressed acoustic-driven activity in the IC. The amount of suppression depended on a number of factors, including the relative time between the acoustic and electrical stimuli, the best
frequency of the A1 and IC electrode sites, and the location of the IC site along an isofrequency lamina. Stimulation of deeper A2 output layers elicited a variety of interesting suppressive results, including direct suppression from stimulation, residual suppression affecting acoustic-driven spiking after stimulation had ceased, and suppression of spontaneous activity.

**Conclusion**
Cortical stimulation can suppress acoustic-driven and spontaneous neural firing in the IC, demonstrating its potential for the treatment of tinnitus. Further work must be performed to characterize the optimal locations and stimulation strategies for eliminating the phantom percept, including counteracting the hyperactivity and hypersynchrony associated with the tinnitus percept.

This work was supported by NIH NIDCD R03-DC011589, NIH NIDA T32-DA022616, the University of Minnesota Institute for Engineering in Medicine Walter Barnes Lange Memorial Award, and start-up funds from the University of Minnesota (Institute for Translational Neuroscience and College of Science and Engineering).

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### Extensive Areas of the Cortex are Evoked by Stimulation from the Newly Implanted Ear in Children who were Long-Term Unilateral Cochlear Implant Users

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**Background**
We recently showed that the development of cortical responses follows a normal-like trajectory with time-in-sound over long-term unilateral cochlear implant use when the duration of bilateral auditory deprivation in childhood is limited. We know from normal hearing data that the temporal lobe is the principal region of auditory processing in the brain. However, the sources of cortical activation in cochlear implant users remain unknown. In the present study, we investigate the areas of the brain that are activated by sound in cochlear implant users. Specifically, we ask whether cortical areas of the brain which normally respond to sound are activated by auditory input in adolescents who grow up using one cochlear implant; and whether activity in the deprived pathways is compromised by the long-term use of only one implant.

**Methods**
Electrically-evoked cortical responses were recorded within the first week of bilateral cochlear implant activation in 12 adolescents who had over 10 years of unilateral cochlear implant experience at the time of the test. Responses were recorded from 64-cephalic electrodes, using 250Hz biphasic pulse trains presented at a rate of 1Hz and evoked by electrical pulses delivered from cochlear implants on the experienced and naïve sides separately. Cortical activation underlying the peaks of the evoked cortical response was localized to different areas of the brain (i.e., temporal, frontal, parietal and occipital lobes) on both hemispheres (ipsilateral and contralateral to stimulation), using our unique beamforming method.

**Results**
Consistent with previous reports, responses from the experienced ear contained amplitude peaks similar to those expected in same aged peers with normal hearing. These peaks were found to come predominantly from the temporal (i.e., auditory) cortex. By contrast, responses from the naïve ear had completely different peaks occurring with larger amplitudes. Neural activity underlying these peaks was more widely distributed, occurring in temporal, frontal, parietal and occipital areas.

**Conclusion**
Long-term unilateral cochlear implant use promotes maturation of the auditory cortex in children. However, diffuse activation of the temporo-fronto-parietal regions from stimulation of the naïve side suggests that the deprived auditory pathways may recruit more attentional resources to process the novel input.
Auditory Cortex Structure and Metabolite Findings in Older Adults with High-frequency Hearing Loss
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Background
Age-related differences in auditory function in older adults with normal hearing may be attributed to central auditory system declines and to changes in the auditory periphery not reflected in pure-tone thresholds. Although thresholds for these individuals may be equivalent to normal at lower and mid-frequencies, most older adults exhibit varying degrees of higher frequency hearing loss. The effects of hearing loss and the sensitivity of pure tone thresholds as they relate to changes in the structures of the central auditory system remain unclear.

Methods
To better understand the effects of high frequency hearing loss on the aging central auditory system, we examined the extent to which variation in high frequency hearing was associated with variation in auditory cortex structure and metabolism in 15 older adults whose pure-tone thresholds were ≤ 25 dB HL from 0.5 kHz to 3.0 kHz and ranged from 15 to 70 dB HL at 8.0 kHz. Diffusion imaging (DTI), magnetic resonance spectroscopy (H-MRS) (NAA), and T1-weighted images were compared to thresholds at 8.0 kHz, where individual differences were largest. Right and left Heschl’s gyri were selected as regions of interest (ROI).

Results
Pure-tone thresholds at 8.0 kHz were positively correlated with DTI mean diffusivity (MD) in the right HG, suggesting that changes in high frequency hearing occur with changes in white matter microstructure. Similarly, elevated thresholds at 8.0 kHz occurred with lower NAA values, suggesting reduced neuronal viability with high frequency hearing loss. Results from linear regression suggest that the association between high-frequency hearing loss, MD, and NAA remained significant after controlling for differences in brain volume and age.

Conclusion
Our measurements were obtained from large ROIs, which encompass regions associated with encoding of both low and high frequency stimuli, and did not have the spatial resolution of voxel-based analyses in which gray matter associations with pronounced high frequency hearing loss have been observed. These results suggest that (1) diffusion and H-MRS metrics are sensitive to auditory cortex changes that may occur with age-related declines in high frequency hearing and (2) widespread central auditory system changes may occur with age-related changes to the auditory periphery that do not result in hearing loss typically considered clinically significant. These changes in white matter integrity and metabolism may contribute to suprathreshold processing deficits observed in older adults with normal hearing.

Supported by NIH

Corticofugal Slow Effects of Auditory Cortex Electrical Microstimulation on Cochlear Responses in the Chinchilla
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Background
The auditory efferent system comprises descending pathways from the auditory cortex to the cochlea, including the cortico-collicular and the olivocochlear system. The electrical stimulation of the medial olivocochlear fibers produces two types of effects with different time scales: (i) fast (tens of milliseconds) and (ii) slow effects (tens of seconds). Whether these two types of effects could also be obtained by electrical stimulation of the auditory cortex is unknown. Here, we recorded cochlear microphonics (CM) and auditory-nerve compound action potentials (CAP) previous, during and after auditory cortex microstimulation and found slow but not fast corticofugal modulations.

Methods
CAP and CM responses were recorded using a right round-window electrode. Cortical evoked potentials were obtained from a Nichrome® electrode positioned in the left auditory cortex of nine anesthetized chinchillas (ketamine/xylazine). Trains of four electrical pulses (0.25 ms each, separated by 2.2 ms) were delivered to the auditory cortex at different rates (1-33 Hz) and current intensities (1-100 µA), during one to five minutes, using an isolated pulse generator (2100, AM-Systems®). Tone bursts were presented at different frequencies (0.5-8 kHz) and intensities (20-90 dB SPL) using a
Tucker-Davis-Technologies® system III hardware and ER-2 (Etymotic Research®). Data was acquired using a National Instruments® Board (NI-6071E).

Results
Auditory-cortex microstimulation produced significant changes in the amplitude of CAP and CM in five experiments. Of these, three (two) showed decrements (increments) of both potentials, 2.3 (2.7) and 2.6 (0.9) dB on average for CM and CAP, respectively. Only one showed modulations going in opposite directions, 1.5 and -8.6 dB for CM and CAP, respectively. These corticofugal modulations appeared only after three to five minutes of electrical stimulation. In addition, these effects were obtained only using microstimulation rates higher than 30 Hz.

Conclusion
Different to olivocochlear electrical stimulation, auditory-cortex electrical microstimulation in the chinchilla produced only a slow modulation of cochlear responses that builds up after three to five minutes. These results suggest that fast olivocochlear modulations are restricted to brainstem circuits, while slow efferent modulations are present from the auditory cortex to the cochlea.

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Functional reorganization in area DZ of congenitally deaf cats
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Background
Sensory deprivation during development can induce functional reorganization of cortical sensory areas. Areas deprived from their natural sensory input may take part in processing of other inputs. In congenitally deaf cats - a model of sensory deprivation – possible visual reorganization of the auditory dorsal zone (DZ) has been demonstrated indirectly in behavioral studies (Lomber et al., Nat Neurosci 2010).

Methods
Here we studied extracellular unit activity in the auditory area DZ and the adjacent visual areas AMLS/PMLS of cochlear-implanted congenitally deaf cats (n=4) and hearing controls (n=4). We measured neuronal responses to visual, electrical auditory and bimodal stimulation simultaneously in AMLS/PMLS and DZ using two 16-channel linear multi electrode arrays (MEA). Recording positions were reconstructed histologically after the experiment.

Results
Modality, distribution and relative frequencies of sensory responsive sites were generally similar in hearing and congenitally deaf cats, in area DZ as well as AMLS/PMLS. Although visual responsive sites in area DZ were sparsely distributed in both groups, the total number of visually modulated sites was increased in congenitally deaf cats.

Conclusion
The results demonstrate that area DZ of congenitally deaf cats remains responsive to input from the cochlea, however, also shows that area DZ of congenitally deaf cats receives increased visual input.
Cortical auditory processing in individuals with long-term symptoms following mild TBI
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Background
While many individuals recover rapidly from concussion or mild traumatic brain injury (mTBI), a significant number of these individuals continue to experience debilitating problems for months and years after injury. Auditory processing problems, including disproportionate difficulty understanding speech in competing noise, are among these common long-term problems. The goals of the current study were to examine neural correlates of auditory processing in this population at both sensory/obligatory and cognitive levels of the central auditory system using cortical auditory evoked potentials and to compare these measures with functional performance on behavioral auditory processing and cognitive test batteries.

Methods
Electrophysiological measures were conducted in a group of individuals who sought rehabilitation in a concussion management program due to ongoing symptoms 3-18 months post mTBI (n =21) and un-injured age and gender-matched controls. Both obligatory cortical auditory evoked potentials (P1-N1-P2) and cognitive event-related potentials (P300) were recorded to speech syllables in quiet and in a background of multi-talker babble. A neuropsychological cognitive test battery and a battery of behavioral tests of central auditory processing were also completed.

Results
Preliminary results showed that over 50% of the mild TBI participants performed abnormally on 2 or more behavioral tests of central auditory processing. Overall, there were few significant group differences in peak latency or amplitude for the P1-N1-P2 or the P300, although P300 behavioral reaction time was significantly prolonged and percent correct discrimination significantly poorer in noise for the mTBI group compared to controls. Within the mTBI group, P1-N1-P2 responses tended to differ more from controls in those who performed abnormally on auditory processing tests but had normal performance on most cognitive tests, while P300 responses tended to differ from controls more in those who performed poorly on both auditory processing and cognitive tests.

Conclusion
A significant number of individuals still symptomatic in the long-term following mTBI showed abnormal performance on behavioral tests of central auditory processing. As a group, there was evidence of slowed processing and reduced discrimination in background noise. Patterns of results from cortical auditory evoked potential and behavioral tests may suggest primarily sensory level auditory processing deficits following mTBI in some individuals, while others have evidence of more global cognitive processing problems that include difficulty processing auditory information. These results could have important implications for rehabilitation planning.
This research was supported by NIH R03DC010246.

Development of a Rodent Model of Chronic Stress-induced Auditory Dysfunction
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Background
Tinnitus and hearing loss are two of the most prevalent auditory impairments. Recent studies have correlated the presence of hearing loss and tinnitus with high levels of life stress. Importantly, the relationship between stress and auditory dysfunction appears multifaceted; not only does hearing loss and tinnitus increase stress, but stress itself may contribute to the development of these auditory disorders. At present, the underlying mechanisms by which chronic stress contributes to auditory dysfunction are unknown; however, chronic stress has been shown to alter GABAergic neurotransmission in the brain. Stress-induced alterations in GABAergic neurotransmission along the auditory pathway represents one possible route by which chronic stress could contribute to auditory disorders. In order to investigate this possibility, we developed a rodent model to characterize changes in auditory brain regions following chronic stress and hearing loss.

Methods
Male Sprague Dawley rats were divided into 4 groups: (1) restraint stress (6 hours/day for 21 days), (2) noise exposure (1h,16-20kHz, 123dB, bilateral), (3) chronic stress plus noise exposure, and (4) control. Tone-evoked auditory brainstem responses (ABRs) were performed at baseline and following stress and noise exposure. Restraint stress was confirmed by monitoring changes in body weight as well as performance on elevated plus maze and open field tests. Standard
immunohistochemical techniques are being used to profile the changes in expression of the GABAergic marker GAD67 in the brain.

**Results**

As predicted, chronic restraint stress resulted in reduced body weight, as well as less time spent in the open arms of the elevated plus maze and in the center of the open field. Restraint stress alone did not affect ABR thresholds. Noise exposure alone significantly elevated ABR thresholds; however, ABR thresholds of rats exposed to chronic restraint plus noise were not significantly different from rats exposed to noise alone. Assessment of changes in expression of GABAergic markers is in progress.

**Conclusion**

Our rodent model will allow us to investigate the underlying mechanisms by which chronic stress contributes to auditory dysfunction. Understanding the contribution of stress to auditory dysfunction may help identify individuals at increased risk for developing hearing loss and/or tinnitus and may lead to novel treatment and prevention strategies.

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**Acoustically-Evoked Auditory Change Complex in Children with Auditory Neuropathy Spectrum Disorder: A Potential Objective Tool for Identifying Cochlear Implant Candidates**

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**Background**

Children with Auditory Neuropathy Spectrum Disorder (ANSD) present a challenge for early intervention and habilitation because of the diversity of the disorder and the current lack of robust indicators to guide in management of this population. This project investigates the utility of the acoustic change complex (ACC) for predicting hearing aid success in ANSD children. It tests the hypothesis that speech perception performance of ANSD children fit with hearing aids can be predicted using the ACC elicited by stimuli containing temporal gaps.

**Methods**

To date, eight ANSD children ranging in age between 1.9 to 14.2 years have been tested. These ANSD children had mild to moderately severe hearing loss between 250-2000 Hz. All subjects had a minimum of 12 months experience with their hearing aids. The stimulus was an 800-ms Gaussian noise presented at 35 dB SL re. pure tone average through ER-3A insert earphones. Two stimulation conditions were tested: (1) In the standard condition, an 800-ms Gaussian noise was presented to the test ear without any interruption; (2) In the gap condition, a silent period (i.e. temporal gap) was inserted after 400 ms of stimulation. The gap duration was fixed at 5, 10, 20, 50, or 100 ms. The ACC was differentially recorded from high forehead (Fz, active) to contralateral mastoid (A1/2, reference) relative to body ground at low forehead (Fpz) . For each gap duration, two replicates of 100 artifact-free sweeps were recorded. These replicates were averaged together for data analysis. Presence of an ACC response required two criteria to be met: (1) identification of a repeatable response complex within the expected latency window; and (2) a root mean square (RMS) amplitude within this window that was at least 50% higher than the noise floor. The shortest gap that could reliably evoke the ACC response was defined as the gap detection threshold.

**Results**

Robust onset cortical auditory evoked responses were recorded from all ANSD subjects. ACC responses elicited by gap stimuli were also recorded from all ANSD subjects. However, subjects who exhibited limited benefit from their hearing aids had longer gap detection thresholds than subjects who received substantial benefit from their listening devices.

**Conclusion**

These preliminary data suggest that the ACC elicited by temporally interrupted stimuli might be predictive of hearing aid outcomes. If so, this objective measure could play an important role in the assessment of ANSD children for cochlear implant candidature.
Automated Fibertracking of the Acoustic Radiation Tract in Normal and Hearing-Impaired Adults
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Background
The acoustic radiation (AR) tract is the penultimate stage of the auditory pathway. It consists of white matter fibers coursing from the medial geniculate body (MGB) of the thalamus to the Heschl’s gyrus (HG) which contains the primary auditory cortex. However the AR presents a challenge to automated methods for tracking white matter fibers in diffusion tensor images (DTI) as it is partly obscured by the optic radiation tract running from the lateral geniculate body (LGB) of the thalamus to the visual cortex. Only probabilistic tracking methods can deal with crossing fibers. One such method is dynamic programming (DP) tracking of single fibers between specified regions of interest (ROIs) and is shown here to reliably generate AR from DTI scans.

Methods
DP uses a quadratic cost based on the Gaussian model of diffusion at the DTI voxel. The HG and MGB were parcellated automatically via the LDDMM registration algorithm in MRI Studio. With the HG and MGB respectively as the start and terminal ROI, DP was used to generate the AR in the left hemisphere in DTI scans of 5 normal and 5 hearing-impaired (HI) subjects.

Results
3D visualization reveals that the fibers are initially bunched together and then begin to spread out like a fan as the AR approaches the HG. Further there is little difference in the fractional anisotropy and spatial topology of the AR in normal and HI subjects.

Conclusion
While DP requires accurate parcellation of ROIs, it is less computationally intensive than other probabilistic tracking methods. The spreading pattern of AR appears to be consistent with published post-mortem staining at least in the coronal view and suggests a structural basis for the HG to be tonotopic. The macroscopic structural integrity of the AR appears to be maintained in HI subjects due to early intervention and may be a contributing factor in positive outcomes in the 4 subjects who subsequently got a cochlear implant. Generation of additional tracts in both hemispheres from the MGB and non-MGB parts of the thalamus to the whole superior temporal gyrus as well as from the LGB to the visual cortex will also be presented. Research supported by P41-EB015909, R01-AG020012 and National Organisation for Hearing Research Foundation.
Nerve survival in CI patients, and its functional consequences: psychophysical measures and cortical recordings

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Background
The objective of the present study is to examine cochlear implant (CI) patients’ sensitivity to pulse phase duration (PPD) as a potential index of nerve survival. We take two approaches to the question: In one approach (Chatterjee lab) we are measuring i) detection threshold for different PPDs and ii) supra-threshold sensitivity to PPD modulation with bipolar (BP), tripolar (TP) and monopolar (MP) stimuli.

In the second approach (Friesen lab) we are conducting parallel measurements of supra-threshold psychophysical PPD discrimination and CAEP responses to changes in PPD, in patients who have had a “softer” surgical implantation with the MEDEL flex soft electrode and those who were implanted with the same electrode but did not have the soft surgery. We hypothesize that the soft surgery will result in improved nerve survival and improved sensitivity to PPD changes in both sets of measures, particularly in apical cochlear regions.

Methods
All stimuli are delivered using custom research interfaces. Detection and discrimination thresholds are measured using forced choice adaptive procedures. We are recording the N1-P2 and ACC on the same CI electrodes for both the low and high range of phase durations to determine the relationship to the behavioral jnd.

Results
Consistent with previous results, results suggest that MP stimulation results in more consistent, efficient charge integration (~ 6 dB/doubling) at threshold than BP stimulation. Strong level-dependence in the across-site pattern of sensitivity to PPD modulation was observed with focused stimuli.

Generally, results show reduced discrimination JNDs and ACC latencies for the longer PPDs. Discrimination JNDs are also smaller for the shorter PPD on the apical electrodes compared to basal electrodes. In the individuals with the “soft” surgery, JNDs are smaller for shorter PPDs on the apical electrodes compared to those not having the soft surgery; reduced ACC latencies were observed for these shorter PPDs on the apical electrodes as well.

Conclusion
Results are consistent with greater dominance of small-diameter peripheral processes in the response, at least at threshold. Sensitivity to PPD modulation likely reveals an increased contribution of central axons at increasing current levels.

With the second approach, it appears that for individuals having the “soft” surgery, PPD JNDs for short duration PPDs are smaller on the apical electrodes, indicating better neural survival. This increased neural survival appears to lead to increased neural synchrony, reflected at the level of the auditory cortex as indicated by the shorter ACC latencies.
Resting cortical oscillations predict hyperacusis: a hyperresponsiveness network with inactive auditory cortex at rest

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Background
Hyperacusis is a hyperresponsiveness to non-noxious auditory stimuli, analogous to allodynia in neuropathic pain, but the neural substrates of hyperacusis are not well understood. By comparing tinnitus participants with hyperacusis (T+H+ group) with those without hyperacusis (T+H- group), we aimed to explore characteristic resting-state cortical activity of hyperacusis.

Methods
The T+H+ and T+H- groups were comprised of 17 participants each and were strictly matched for all tinnitus characteristics (tinnitus intensity and distress, laterality and type, gender, onset age and duration) to exclude cortical activity changes due to tinnitus characteristics. We compared differences in cortical activity between the 2 groups using resting state electroencephalography (EEG) source-localized activity complemented by functional connectivity analyses. Also, EEGs of 17 normal controls, matched for sex and age, were utilized for further comparison.

Results
Compared to the T+H- group, the T+H+ group demonstrated increased beta power in the dorsal anterior cingulate cortex (dACC), orbitofrontal cortex (OFC) and supplementary motor area (SMA) and increased alpha power in the right auditory cortex (AC). Region of interest analyses including normal controls further confirmed that these differences originated solely from relatively increased power of the T+H+ group, while power was not increased relative to the normal hearing controls. Functional connectivity analysis revealed increased connectivity between the OFC/dACC and the AC in the T+H+ group in comparison to the T+H- group. The beta power increase in the OFC/dACC/SMA may indicate increased vigilance leading to increased startle reflex in hyperacusis participants. Additionally, increased resting-state alpha power in the AC may reflect an adaptive top-down inhibition against environmental sound stimuli probably mediated by the increased beta power of the OFC.

Conclusion
The OFC and the dACC may compose a hyperresponsiveness network as these areas have also frequently been found to be activated in allodynia/hyperalgesia studies, and the results may be applicable for developing future treatment strategy in patients with hyperacusis or allodynia/hyperalgesia.

The Role of GABAergic and Glycinergic Inputs in Shaping SSA in the Inferior Colliculus of the Anesthetized Rat.

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Background
Stimulus-specific adaptation (SSA) is a reduction in the responsiveness of a neuron to a common or repetitive sound while the neuron remains highly sensitive to a deviant sound (Ulanovsky et al., 2003). This single-neuron level phenomenon could enhance the saliency of unexpected stimuli against a background of repetitive signals. Previous work from our laboratory has shown that adaptation to the repetitive stimulus still occurred in the absence of GABAA function and that the GABAA-mediated inhibition acts as a gain control mechanism that enhances SSA by controlling the neuron’s gain and responsiveness (Pérez-González et al., 2012). Here, we test whether inhibitory mechanisms other than those GABAA-mediated contribute to the generation and/or modulation of SSA in the inferior colliculus (IC).

Methods
Antagonists of GABA_A+, GABA_B- and glycinergic receptors (gabazine, CGP-35348 and strychnine, respectively) were applied microiontophoretically while recording single units in the IC of rats, using an oddball paradigm. The responses to high- and low-probability tones were recorded before, during and after the drug injection using an assembly of microelectrodes in a ‘piggy-back’ configuration. The experimental stimuli were pure tones in the range 0.5–40 kHz, with a
Results
So far, we have recorded the extracellular activity of 30 single units. The application of gabazine and CGP-35348 increased spontaneous activity and altered the tuning of the FRAs and the rate-level functions at the best frequency, although the magnitude of the effect was higher under gabazine. The blockade of the three different receptors independently increased the response magnitude to high and low probability stimuli and caused a reduction of the first spike latency. Application of CGP-35348 alone had only a weak effect on SSA. However, when combined with gabazine and/or strychnine the effects were more profound than each drug alone and resulted in a significant reduction of SSA.

Conclusion
Our data indicates a synergic action of GABAergic and glycinergic inhibition in shaping SSA at the IC level. Furthermore, it suggests that the SSA observed in the IC is generated locally.

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Response to Complex Patterns of Regularity in the Inferior Colliculus of the Anesthetized Rat
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Background
Single neurons in the inferior colliculus (IC) of the rat show a specific decrement of response to simple repetitive stimuli. This ability to encode the complex auditory past in multiple time scales would enable the auditory system to generate expectations about the incoming stimuli. So far, stimulus-specific adaptation (SSA) has been studied in animal models using the simple repetition suppression effect of the classical oddball paradigm. Yet it remains to be determined whether violations of more complex regularities can also be detected at single neuron level in the form of expectation suppression, as they are in humans at population level through EEG and MEG.

Methods
Experiments were performed on female Long-Evans rats anesthetized with urethane (1.5 g/kg, i.p.). Tungsten electrodes (1–2 MΩ) were used to make extracellular recordings of neural responses in well-isolated single units in the IC. Auditory stimuli were sequences of 75 ms long pure tones with a 5 ms rise/fall ramp and constant stimulus-onset asynchrony (SOA) of 250 ms. Sound sequences consisted of two alternating pure tones selected from the frequency response area (FRA) of the neuron, following an alternation-repetition paradigm similar to those used in human studies. The average response to each stimulus when it was embedded in a regular pattern (expected), was compared to the average response to that same stimulus when it was breaking the regularity (unexpected). We presented each stimulus as repeated-expected and repeated-unexpected in order to control for repetition suppression, and we tested omission trials as well to look for true pattern regularity violation sensitivity.

Results
Among the recorded neurons, a significant number of them were considered strongly adapted neurons, showing much higher response to deviant tones than to the same tones being standard within a classical oddball paradigm. By contrast, other units showed little or no SSA, i.e., similar response profiles to standard and deviant stimuli. Remarkably, we have found signs of pattern violation sensitivity in both classes of neurons, in the form of a differential response to expected and unexpected tones, regardless of its repetition.

Conclusion
Our preliminary results show that at the level of the midbrain, there is not only repetition suppression (SSA) but may be also expectation suppression. These properties may occur independently to a different extent in different neurons, but this question awaits future experiments.

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Stimulus Specific Adaptation in the Inferior Colliculus of the Awake Marmoset
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Background
Neural mechanisms to detect novel stimuli in perceptually complex environments are fundamental to our survival. In sensory systems, these mechanisms exist at a cellular level in the phenomenon known as stimulus specific adaptation (SSA) where single neurons can be seen to adapt to repetitive (standard) stimuli while remaining highly responsive to rare stimuli. While SSA has been observed in the auditory system at the level of the cortex, thalamus and the inferior colliculus, there remains debate as to its origin particularly given the paucity of data from unanaesthetized preparations with unmodified inhibitory circuits.

Methods
To further elucidate this system, single neurons were recorded from the inferior colliculi of awake head-fixed marmoset monkeys with tungsten microelectrodes. SSA was assessed with five free-field presentations of three consecutive streams containing repeated tones of two frequencies, f1 and f2 chosen from either side of the best frequency of each neuron. In the first stream, f1 was presented with high probability (90%), and f2 at low probability (10%), these probabilities were inverted in the second stream and presented at equal probability in the third stream.

Results
The majority of neurons within the central (ICC) and external (ICX) nuclei showed some degree of SSA. These effects were observed in both sustained and onset neurons although the latter were less frequently observed. This phenomenon was frequency specific such that SSA was greatly affected by the position of the tone within the receptive field of the neuron. Indeed latency was also observed to vary throughout the stream in a manner dependent on frequency. This phenomenon was most striking in a number of ICX units that showed a progressive increase in latency over the duration of the stream with a concomitantly large degree of adaptation that did not affect the deviant.

Conclusion
The majority of units recorded with SSA demonstrated sustained discharges throughout tone presentation while some of these showed pauser or attenuated responses after the first 10-15ms. These data convincingly demonstrate that SSA is seen in the central nucleus of the unanaesthetized IC albeit at reduced levels relative to the external nucleus. Many presentation frequencies delivered to ICX units with broad receptive fields induced progressively increased latencies over each stream in a manner dependent on the position of this frequency in the receptive field of the neuron.

Spectral receptive field plasticity may underlie stimulus-specific adaptation in rat inferior colliculus
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Background
Neuron can change its coding strategy dynamically according to its stimulus context. One example in auditory system is stimulus-specific adaptation (SSA), in which rare stimuli elicit higher responses than common stimuli. SSA to frequency was observed at single neuron level in both cortical (Ulanovsky et al., 2003) and subcortical areas (Malmierca et al., 2009; Zhao et al., 2011). However, SSA can only reveal the adaptive change of responses to two specific frequency probes. It remains unknown how frequency selectivity changes in repetitive context and how much these changes relate to SSA. We explored this question by comparing frequency tuning curves of neurons in rat inferior colliculus (IC) in an adapted condition and a normal condition.

Methods
Normal frequency tuning curves were measured at random frequency points centered at neuron’s best frequency (BF) while adapted tuning curve was measured by the same set of probes except they were randomly interspersed in a repeating tone sequence with a fixed frequency. The repeating tones acted as common stimuli and accounted for 90% of the tone sequence. The frequency of the repeating stimuli was selected similar to the standard frequency in the oddball paradigm (Ulanovsky et al., 2003; Zhao et al., 2011).

Results
We found that in adapted condition, the responses to frequencies very close to the repeated one were significantly decreased while to frequencies at farther vicinities showed enhancement. The suppressed/enhanced pattern was
frequency specific with radically center-surround pattern relative to the repeating frequency and depended on both inter-stimulus interval (ISI) and the segregation between the common stimuli and BF of the neuron. Moreover, we stimulated neurons using oddball paradigm with the same common stimuli and found the responses to rare ones can be largely predicted by the adapted tuning curve. Finally, a channel-based adaptation model was proposed, which can qualitatively predict the observed change in adapted condition. It revealed that adaptation induced frequency specific suppression of input to IC through the underlying center-surround connection pattern inter- and intra- IC neurons.

Conclusion
This result suggests that neurons’ receptive fields undergo elaborate dynamics during adaptation to the ongoing sound stream, which may increase the frequency discriminability to facilitate novelty detection. Furthermore, it provided evidence of the neural mechanism of SSA and suggested more properties of adaptation, which were not shown by current SSA experiments.

Temporal Modulation Tuning in the Auditory Midbrain Provides Distributed Neural Representation for Coding Vowel-like Sounds
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Background
Many auditory midbrain neurons are tuned to amplitude modulation frequencies between 50 to 500 Hz. It is reasonable to ask whether these neurons play a particular role in coding speech information and if so, how? It has been hypothesized that midbrain neurons that are periodicity-tuned to voice pitch will have decreased rates when the neuron’s best frequency (BF) is near a formant frequency and increased rates when BF is between formants (Carney and McDonough, 2012, JASA 131:3309). The present study tests this hypothesis across a wide range of best frequencies using synthesized vowel-like sounds to explore the role of temporal modulation tuning of neurons in speech coding.

Methods
Responses of single neurons in the inferior colliculus (IC) were recorded in the awake rabbit. Response maps and modulation transfer functions (MTFs) were measured to characterize neurons. Stimuli were presented to the contralateral ear. The stimuli were synthesized harmonic complexes with vowel-like spectra having two formants. The two formants were shifted with respect to the BF of the neuron to test the above-mentioned hypothesis. The fundamental frequency of the stimulus was matched to the best modulation frequency of the neuron. Two phase relationships (random and sine phase) across the harmonics were used to generate two sets of stimuli with similar spectra but different temporal envelopes. The average discharge rate and synchronization coefficient were used to quantify the neural responses.

Results
For the neurons with narrowband MTFs the responses to the vowel-like harmonic complexes with sine phase were significantly stronger and more synchronized to the stimulus than responses to random phase complexes. For onset neurons with broad MTFs, the responses to the vowel-like harmonics complexes with sine phase were also significantly stronger and synchronized to the stimulus. For sustained neurons with broad MTFs, the difference in the responses to the two sets of stimuli diminished. Some neurons had decreased discharge rates when a simulated formant was near the BF of the neuron, supporting the formant-coding hypothesis.

Conclusion
The present study provides evidence to support the hypothesis for coding vowels at the level of the auditory midbrain. The results also show that IC neurons with different modulation transfer functions may encode stimuli with similar spectra but different temporal envelopes in different manners, suggesting a distributed neural representation for temporal information in speech in the auditory midbrain.

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Serotonin Reduces Noise-evoked Suppression of Vocalization Responses in Inferior Colliculus
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Background
One of the ways the neuromodulator serotonin influences the excitatory-inhibitory circuitry of the inferior colliculus (IC) is by decreasing the strength of the GABAergic inhibitory surround.

Methods
To explore a functional consequence of this interaction in CBA/J mice, we used a stimulus consisting of a vocalization alone or one paired with a broadband noise. For a given neuron, vocalizations eliciting a strong excitatory response were chosen from a range of sample vocalizations and played at levels from just below threshold to 20 dB above threshold to generate a limited rate-intensity function. Noise was presented at a level for each neuron that evoked a minimal response alone while influencing the response to the vocalization.

Results
Many neurons in the sample showed effects similar to those previously reported in the IC for the influence of noise on tone responses, including an increase in threshold and a shift in the rate-intensity function to higher intensities. Noise suppressed the spike counts in response to vocalizations near threshold by an average of 41%. Serotonin reduced the effect of noise, so that noise only decreased the spike count by 25% on average in the presence of serotonin in the same group of neurons. However, neurons varied widely; serotonin almost entirely removed the effects of noise in some, but even facilitated the effects of noise in others. Noise also altered the timing of spike trains in some neurons, an effect that could be abolished by serotonin.

Conclusion
These results are consistent with multiple mechanisms of serotonergic action, including a decrease in the surround inhibition leading to a reduced influence of broadband noise for a limited range of vocalization-noise intensity pairs.
Responses to Acoustic Stimuli Vary Along the Isofrequency Laminae of the Inferior Colliculus Consistent with Spatially Segregated Sub-lemniscal Auditory Pathways
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Background
The central auditory system has traditionally been divided into the lemniscal and non-lemniscal pathways. The lemniscal pathway transmits high-fidelity auditory information and maintains a precise tonotopic organization. In contrast, the non-lemniscal pathway codes for multi-modal information and is involved with modulation of the lemniscal pathway and limbic interactions. The central nucleus of the inferior colliculus (ICC) is the main auditory midbrain nucleus of the lemniscal system. Based on previous anatomical and neurophysiological studies, we hypothesize that the ICC should be further divided into at least two sub-regions along the isofrequency laminae, a rostral and a caudal region, that exhibit different coding properties to sound.

Methods
Short tone pips were presented to ketamine-anesthetized guinea pigs. Multi-site electrode arrays were used to record local field potentials (LFPs) and spikes simultaneously in up to four regions along an ICC lamina. Multiple array placements were made for each animal. Three-dimensional histological reconstructions of the placements were performed for six animals to identify all the recording locations across a given ICC lamina. The responses were then compared across locations.

Results
We observed that both LFP and spike response patterns differed between rostral versus caudal regions along an ICC lamina. The rostral region exhibited more temporally-precise spiking, shorter first spike latencies, less jitter in first spike latencies, and shorter response durations. The rostral region also exhibited LFPs with shorter peak latencies as well as activity with greater growth and temporal synchrony.

Conclusion
These results support previous histological and electrophysiological studies suggesting that the lemniscal pathway should be divided into at least two sub-projection pathways. Specifically, the rostral region appears to have properties more consistent with the lemniscal pathway (e.g., excitatory discharge patterns, short latencies, and precise time-locking) and may serve to reliably and synchronously transmit sound information from the auditory nerve up to the auditory cortex. Based on additional studies from our lab, the caudal pathway may be more involved with modulating sound information ascending through the lemniscal pathway. Identifying the spatial coding pattern across the isofrequency laminae of the ICC has major implications not only for understanding central auditory processing but also for improving auditory midbrain implants, in which the rostral area of the ICC may be a better stimulation target for restoring intelligible hearing. This research was funded by University of Minnesota start-up funds and NIH NIDCD R03DC011589.

Progressive Recovery of Acoustic Sensitivity in the Inferior Colliculus of Adult Mice Following Near-Complete Primary Cochlear Neuronal Degeneration
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Background
Spiral ganglion neuron (SGN) survival declines as a normal consequence of aging and in pathological conditions such as auditory neuropathy spectrum disorder and acoustic trauma.

Methods
To better understand how SGN loss affects the central auditory system, we performed electrophysiological recordings from the central nucleus of the inferior colliculus (ICC) of adult mice following specific depletion of Type I SGNs in the contralateral cochlea. For each individual mouse, in vivo multunit recordings were compared to the auditory brainstem response (ABR), distortion product otoacoustic emissions (DPOAE), histological quantification of inner hair cell synaptic complexes or the spiral ganglion, and acoustic startle reflex behavior.

Results
Unilateral application of ouabain to the round window membrane eliminated $\sim$90\% of Type I SGN synapses onto inner hair cells without affecting hair cell function or Type II SGN survival, as determined by quantitative histopathology and
otoacoustic emission measurements. As expected, the amplitudes of the auditory brainstem response and acoustic startle reflex elicited from the ouabain-treated ear were substantially reduced. Despite the profound deafferentation and apparent hearing loss, a substantial minority of ICC units exhibited low-threshold, non-adapting, and tonotopically organized sound-evoked responses one month – but not one week – following ouabain treatment. By contrast to the recovery of responses to simple sounds such as tones in silence, the representations of tones presented in broadband masking noise or the accurate synchronization of spike times to the modulation rate of the sound pressure envelope were similarly impaired at both recovery times.

Conclusion
The persistence of robust responses evoked from the ouabain-treated ear could not be attributed to the fraction of surviving inner hair cell synapses, but rather appeared to represent a slowly developing compensatory plasticity that was evident at the level of midbrain unit responses, but not in the gross electrical potential or startle reflex generated by brainstem circuitry. These findings imply a readjustment of synaptic gain in the ascending auditory pathway, which may have relevance to the development of tinnitus, hyperacusis or other complex behavioral sequelae of peripheral damage.

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Masking Release due to Coherent Envelope Fluctuations across Frequency at the level of the Inferior Colliculus
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Background
Many natural sounds including speech contain coherent level fluctuations in different frequency bands. A psychoacoustical phenomenon associated with the ability of the auditory system to use this characteristic is comodulation masking release (CMR). The physiological mechanisms underlying CMR are still unclear.

Methods
The present study used a typical CMR paradigm, in which a tonal signal is embedded in a masker with a spectral component at the signal frequency and one or more off-frequency components. The psychoacoustical CMR describes the effect of a reduced masking when the masker components showed coherent level fluctuations (comodulated condition, CM) compared to a condition in which the level fluctuations were incoherent (codeviant condition, CD) or to a condition where the masker consisted of the on-frequency component only (the so called reference condition, RF). The present study used sinusoidally modulated tones as masker components, one centred at the best frequency of the unit and the other positioned into the inhibitory sidebands of the unit. The recording was done in the inferior colliculus of the Mongolian gerbil; 53 units in 17 animals were tested.

Results
Preliminary analysis on the current data base showed that 33 out of 53 units were able to follow the amplitude modulation. Out of these ca. 25% displayed response properties that were consistent with the hypothesis of wideband inhibition, the hypothesized physiological mechanism underlying CMR from previous studies at the level of the cochlear nucleus. Approximately 20% of the units also showed a reduction of the representation of the masker modulation due to the presence of the signal, i.e., envelope locking suppression which is another hypothesized physiological mechanism. This mechanism was proposed on the basis of cortical recordings. Since both effects were found at the level of the IC, it is possible that the auditory system uses a combination of both, leading to a further enhancement of signal detectability.

Conclusion
Wideband inhibition is a mechanism that may account for a physiological correlate of CMR at the level of the IC, and additionally envelope locking suppression could also play a role.
Auditory Stream Segregation Abolishes Informational Masking

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Background
Informational masking describes a common phenomenon observed in everyday life. Important acoustic information, like an alarming sound, can be masked by surrounding acoustic events, even up to the point in which the important information is below detection threshold although the sound is well represented in the auditory periphery. This non-energetic masking is thought to originate in the central auditory system and can be measured and defined as an increase in detection threshold caused by stimulus uncertainty (e.g., Durlach et al. 2003). Most studies have concentrated on analyzing the effect of simultaneous informational masking (e.g., target and masker sequences are presented simultaneously, see Kidd et al. 2007). In an EEG study of auditory stream segregation Winkler et al. (2003) used the mismatch negativity (MMN) response to evaluate the ability of the brain to detect a deviant (76 dB SPL) in a series of standards (61 dB SPL) and intervening signals (66, 71, 81 and 86 dB SPL). Large differences in MMN response were observed as a function of frequency difference between the standards/deviants and intervening signals.

Methods
We adopted Winkler’s paradigm and applied it in a psychophysical (humans and gerbils) and neurophysiological approach (extracellular recording in the inferior colliculus (IC) and auditory cortex (AC) of anesthetized gerbils). Psychophysical thresholds were obtained with the method of operant conditioning using positive reinforcement in a Go/NoGo procedure.

Results
Similar to other objective tasks of auditory stream segregation (e.g., Micheyl and Oxenham, 2010) the intensity increment thresholds derived with Winkler’s paradigm, providing sequential informational masking, varied depending on the frequency difference between the tones. In both humans and gerbils thresholds were considerably larger in the condition of small frequency differences representing a 1-stream percept than in the condition of a large frequency differences representing a 2-stream percept. No difference between species was observed. Using the same types of stimuli as in the behavioral study, neuronal rate responses were analyzed using the receiver operating characteristic (ROC) functions and dₐ -values were calculated and compared to the psychophysical results. The observed change in the dₐ -values with stimulation condition could explain the behavioral observations.

Conclusion
The similarity between the neurophysiological and the psychophysical response measures in the gerbil renders it suitable as a model revealing the relation between auditory stream segregation and informational masking that may also apply to human perception.
Neural Discrimination of Vocoded Consonants
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Background
Cochlear implants provide an input to the auditory system that is both spectrally and temporally degraded. Despite this, many cochlear implant users are able to discriminate between speech sounds with a performance comparable to normal hearing listeners. Vocoders have been used to acoustically simulate this degraded input in many psychophysical studies, but the neural bases of this robust phonemic discrimination are not yet clear. Understanding the aspects of the neural representation of speech sounds that are used for discrimination and that are robust to degradation of the input signal could have implications for both cochlear implant design and automatic speech recognition systems.

Methods
A set of 16 medial consonants (a/C/a) produced by 3 male speakers were processed using a noise-band vocoder. The number of noise bands was varied between 1, 2, 4 and 8 and temporal modulation bandwidth was limited to 16 and 500 Hz. Multi and single unit responses to the processed and unprocessed stimuli were recorded in the inferior colliculus of anaesthetised guinea pigs. A nearest neighbour classifier was then trained on the average spatio-temporal neural representation of each consonant. Novel activity patterns were classified according to the closest of the training set given any possible relative onset time.

Results
The classifier was able to correctly identify unprocessed phoneme sequences based on neural activity patterns with a success rate of 93%. For the single channel, 16 Hz envelope bandwidth condition, performance dropped to 51% and increased monotonically with increasing numbers of noise bands to 84% for the eight channel condition. For the 500 Hz envelope bandwidth condition, performance was greater than 90% for any number of noise bands.

Conclusion
When envelope modulation bandwidth is restricted to 16 Hz, the distinguishability of neural activity patterns closely matches human psychophysical data for the same task. However, there is sufficient information encoded in the guinea pig auditory midbrain to facilitate near-perfect discrimination of vocoded vowel-consonant-vowel phoneme sequences when temporal modulations of up to 500 Hz are conserved in the input signal. This suggests that although the auditory midbrain preserves temporal information about speech which can distinguish between different speech sounds, that this is not used for speech recognition. One explanation for this might be that generalized representations built on finer temporal information are not robust across environmental variations, such as inter-speaker differences, background noise or reverberation. However, the cause of this is still being investigated.

Mathematical Modeling of Gap Detection
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Background
As part of a study of hearing loss in HIV-positive individuals, we have collected gap detection thresholds on 350 HIV-positive and HIV-negative individuals. For each subject, at each gap length, the percentage of time the subject correctly identifies the gap is measured.

Methods
The dependence of correct identification rate on gap length is sigmoidal, showing a threshold below which no gaps are detected, a rapid rise in detection percentage as gap length increases, and then a plateau. The dataset was systematically analyzed via curve fitting to provide a simple two-parameter characterization of each subject’s gap response. A simple neural network model for gap-sensing was also developed, incorporating a population of neurons, where each neuron has its own sigmoidal response to gap detection (and different gap detection performance).

Results
The gap detection curves fit extremely well to the Hill equation (see figure), which is often used to describe the cooperative binding of multiple ligands to a macromolecule, such as the binding of oxygen to hemoglobin. The equation provides a Hill coefficient, which captures the degree of cooperativity of the binding reaction (i.e. how much the binding of one ligand is enhanced by the presence of others). The Hill equation also naturally emerges from the simple neural-
network model for gap sensing, enabling us to ascribe meaning to the equation’s two parameters, where “n” or cooperativity, is a measure of the number of neurons, and “K” represents the effective gap detection threshold.

**Conclusion**
The Hill equation provides a simple mathematical description of central nervous system gap detection. This process involves multiple neurons tuned to detect particular gap lengths, with the ensemble response providing the overall gap encoding. The concept of cooperativity lends itself to a particularly intuitive interpretation in this setting, which is explored in this study. We show that the Hill coefficient is directly related to the number of neurons involved in our neural network model, whereas the binding constant from the Hill equation corresponds to the gap detection threshold. Although these findings are preliminary, they indicate that the shape of the percent correct vs. gap length curve may reflect neuronal loss in the gap detection pathways. Cooperativity in gap detection may serve to reduce noise in signal detection, similar to the effects of cooperativity in cellular signal processing.
Neural Correlates of the Detection of Tones in Noise in the Inferior Colliculus of Nonhuman Primates

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Background

Detection of sounds in noise is a fundamental function of the auditory system. The thresholds of macaques for tone detection displayed a characteristic U-shaped relationship with frequency. Reaction times to detect tones decreased as the tone sound pressure levels increased beyond threshold. Noise modified the responses by shifting thresholds to higher sound pressure levels at the rate of 1 dB per dB of noise, but reaction times were not significantly different in the presence of noise. The response magnitude of inferior colliculus (IC) neurons was more strongly related to detection of tone alone than those of cochlear nucleus (CN) neurons. Studies in decerebrate and anesthetized cats have shown that the dynamic range of some IC neurons were shifted at the rate of 1 dB per dB of noise, but most CN units showed smaller shifts. In addition, neurons in decerebrate and anesthetized cats showed a compromised representation of signal. Here we report results of experiments designed to evaluate the role of the neurons in the IC of awake and behaving animals in detecting tones in noise.

Methods

Two macaques (*Macaca mulatta*) were trained in a reaction time lever release task with appropriate catch trials to report the detection of tones in noise. Single units were recorded from the IC (n=70) during behavior. The characteristic frequency of these units ranged up to 25 kHz. Most of the units were classified as type V or type I, based on the distribution of excitation and inhibition in frequency response maps. Signal detection theory and Receiver Operating Characteristic (ROC) analyses were used to create psychometric (behavioral), and neurometric functions. Neurometric functions reveal the probability of correct response based on the distribution of the number of spikes fired by the IC neuron on tone trials and catch trials.

Results

Background noise increased the average firing rate of IC neurons. The dynamic range of the neurometric functions shifted to higher sound pressure levels in the presence of noise. On average, with the addition of noise, IC neurometric thresholds showed shift rates of 1 dB per dB noise, and were strongly correlated with behavioral thresholds in noise. The relationship between reaction times and spike counts was not significantly different with the addition of noise.

Conclusion

These results, that the neuron-behavior correlations were not altered in the presence of noise, suggest an important role for the IC in processing signals in noise. Supported by R01 DC11092.
Glycinergic and GABAergic Input Provide Excitatory Drive in the Inferior Colliculus.
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Background
Glutamate is the dominant excitatory neurotransmitter in the auditory system, but there is growing evidence that glycine can provide the sole excitatory input to inferior colliculus neurons through an excitatory rebound from inhibition. This study of the pallid bat inferior colliculus shows that either glycine or GABA can provide excitatory drive to a population of highly selective neurons that respond only to downward FM sweeps, and whose selectivity is shaped by an asymmetrical facilitation when presented with two tones in a spectrotemporal sequence approximating their occurrence in an excitatory downward FM sweep.

Methods
Five-barrel microelectrodes were used to record single neurons, and iontophoretically apply blockers of AMPA (NBQX) and NMDA (CPP) glutamate, glycine (strychnine) and GABAa (gabazine) receptors. Iontophoretic currents were gradually increased to maximize effect, but minimize response saturation.

Results
As expected, the blockade of glutamate receptors suppressed responses in the majority of neurons, and the blockade of glycine and/or GABAa receptors disinhibited responses and altered response selectivity in terms of FM sweep direction or sweep rate. Neurons that previously responded only to downward sweeps now responded to tones as well. However, in 33% of neurons in which glycine receptors were blocked, their response was eliminated. The same effect was observed in 20% of neurons in which GABAa receptors were blocked. In some of these neurons, the blockade of glutamate receptors had no effect on their response, suggesting these inhibitory transmitters provided the sole excitatory drive.

Conclusion
The excitatory effect of glycine and GABA is thought to be the result of a rebound from inhibition, in some cases in coincidence with excitatory input, and has been observed in avian and mammalian thalamic and cortical circuits. In the mammalian auditory system, the excitatory effect of glycinergic input has been documented in both the midbrain and brainstem, and is thought to be a source of temporally precise interneuron communication. Present results indicate that both GABA and glycine, most likely through inhibitory rebound, can contribute to the submillisecond temporal resolution observed in inferior colliculus neurons in which asymmetrical facilitation shapes their selectivity for FM sweep direction and rate.

Identification of Three Classes of Excitatory Synapses in the Inferior Colliculus
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Background
The inferior colliculus (IC) integrates ascending auditory input from the lower brainstem and descending input from auditory cortex. Understanding how IC cells integrate these inputs requires identification of their synaptic arrangements. Here, we provide the first description of excitatory synapses in the main IC subdivisions (dorsal cortex, central nucleus, and external cortex) in guinea pigs. We describe characteristics that allow distinction between excitatory and inhibitory synapses as well as differentiation of 3 classes of excitatory boutons.

Methods
We used electron microscopy (EM) on aldehyde-fixed tissue from 6 pigmented adult guinea pigs. We stained the ultrathin sections with post-embedding anti-GABA immunogoldhistochemistry to identify GABAergic cells. We analyzed 450 synapses (150 from each IC subdivision). Excitatory synapses were identified by round vesicles, asymmetric synaptic junctions, and GABA-immunonegative presynaptic boutons. Presumptive inhibitory synapses had non-round vesicles, symmetric junctions, and GABA-positive (GABA+) or GABA-negative presynaptic boutons.

Results
Excitatory synapses constitute ~ 60% of the synapses in each IC subdivision. Three subtypes can be distinguished by presynaptic profile area and number of mitochondrial profiles. Large excitatory (LE) boutons are more than 2 µm² in area and usually contain 4 or more mitochondria. Small excitatory (SE) boutons are less than 0.7 µm² in area and usually contain 1 or no mitochondria. Medium Excitatory (ME) boutons are intermediate in size and usually contain 2 to 4 mitochondria. A k-means cluster analysis confirmed that the 3 types of boutons could be distinguished by presynaptic
Conclusion
Excitatory boutons give rise to about 60% of the synapses in each IC subdivision. Excitatory boutons comprise 3 subtypes that are distinguishable by profile size and mitochondrial content. SE boutons include boutons that arise from auditory cortex. Most or all LE boutons and many ME boutons likely arise from lower brainstem auditory nuclei. The data suggest that each of these inputs are likely to contact both GABAergic and non-GABAergic IC cells, and thus likely to activate both excitatory and inhibitory IC circuits.

Inhibitory Projections and GABAergic “Cross-talk” between Parallel Pathways in the Projections from Inferior Colliculus to the Medial Geniculate Body in Guinea Pigs.
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Background
Classic studies in cats identified 3 parallel streams in the projections from the inferior colliculus (IC) to the medial geniculate body (MG) (Calford and Aitkin, 1983, J Neurosci 11:2365). The streams are reflected in subdivision-specific connections: 1) a “lemniscal” pathway is formed primarily by projections from central IC to ventral MG (ICc-MGv); 2) a “diffuse” pathway involves projections from IC dorsal cortex to dorsal MG (ICd-MGd); and, 3) a “polysensory” pathway involves projections from IC external cortex to medial MG (ICx-MGm). Smaller numbers of cells in each pathway terminate in the “wrong” MG subdivision, providing an opportunity for interactions, or “cross-talk”, between the parallel pathways. More recent studies have identified a substantial GABAergic projection from the IC to the MG, but whether these projections follow the parallel organization is not known.

Methods
We combined retrograde tracing with immunohistochemistry in guinea pigs to identify GABAergic cells that project to MGv, MGd or MGm. We injected Fast Blue, FluoroGold or red or green fluorescent microspheres into individual MG subdivisions and stained the IC with anti-glutamic acid decarboxylase (GAD) to identify GABAergic cells. We quantified tracer- and immuno-labeled (double-labeled) cells in the IC ipsilateral to the MG injection. The present analysis is based 4 cases (1 for each MG subdivision) and >5,800 retrogradely-labeled cells.

Results
After MGv injection, 58% of the GABAergic tectothalamic cells were in ICc. After MGd injection, 60% of the GABAergic tectothalamic cells were in ICd. After MGm injection, ICx contained 69% of the GABAergic tectothalamic that project to MGm. Thus, the bulk of GABAergic tectothalamic cells are located in IC subdivisions predicted by the classic parallel pathways. However, in each case a substantial percentage of GABAergic tectothalamic cells were located in “other” subdivisions. For example, MGv injections labeled GABAergic cells in ICx and ICd; MGd injections labeled GABAergic cells in ICx; and MGm injections labeled GABAergic cells in ICc.

Conclusion
We conclude that the bulk of GABAergic tectothalamic cells project in a pattern that reflects the classic lemniscal, diffuse and polysensory pathways. However, additional GABAergic IC cells project “across” the pathways, providing an opportunity for inhibitory crosstalk between the parallel pathways in the tectothalamic projection. Supported by NIH DC04391 and DC012450.

c-fos Immunolabeling Reveals Changes in Sound Frequency Representation in Inferior Colliculus Following Chronic Low-Level Neonatal Sound Exposure
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Background
Sensory areas of the brain have the remarkable ability to reorganize as a result of significant changes in peripheral sensory input, especially during early development. While cortex is often investigated as the primary site of developmental
plasticity, our study provides evidence for midbrain plasticity. Our working hypothesis is that the development of auditory pathways is influenced in large part by patterns of sensory activation experienced during early development. Specifically, we hypothesize that post-natal exposure to an unusual sound environment modifies the neural representation of the exposure sound frequency in inferior colliculus (IC).

**Methods**
Newborn chinchillas were reared in the presence of a chronic, moderately-intense (c. 70 dB SPL) narrowband sound signal (centered at 2 kHz) for 4 weeks. We estimated hearing thresholds using auditory brainstem responses (ABRs). We then observed neural activation patterns in IC using c-fos labeling (protocol optimization for chinchilla developed in-house, using commercially-available reagents).

**Results**
Hearing thresholds were similar between control and sound-exposed subjects, suggesting that sound-exposure does not affect normal hearing. We observed a statistically-significant increase in the number of labeled cells both in the 2-kHz region (corresponding to the sound-exposure frequency) as well as throughout the IC of sound-exposed subjects, compared with controls.

**Conclusion**
These results support the hypothesis that abnormal sound patterns at the periphery during a neonatal period can induce changes in neural activation patterns in sub-cortical auditory structures. We also discuss some plausible interpretations. Studying the effect of sound on the developmental plasticity of the auditory brain is important for our general understanding of how the auditory system develops. The work also has some relevance for the rehabilitation of the hearing impaired, for example with hearing aids and cochlear implants.
Reciprocal Connections Between the Inferior Colliculus and Dorsal Cochlear Nucleus

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Background

Descending pathways are an essential element of sensory systems, and may facilitate realtime modification of neural responses to external stimuli at any stage from periphery to cortex. In the auditory system, the cochlear nucleus is a key site for descending influence because it initiates all ascending pathways. Studies of descending projections in rat and guinea pig suggest that colliculo-cochlear nucleus projections originating in the central nucleus of the inferior colliculus (CNIC) terminate bilaterally and topographically in the dorsal cochlear nucleus (DCN; Caicedo & Herbert, 1993; Malmierca et al., 1996). We recently quantified and confirmed this projection in the mouse, and furthermore suggested that this pathway is reciprocal (Muniak et al., 2011). We now present ultrastructural evidence that these descending terminals from the CNIC form excitatory axodendritic and axosomatic synapses on the principal projecting cells of the DCN.

Methods

Adult CBA/CaJ mice received an injection of biotinylated dextran-amine (BDA) and cholera toxin subunit-B (CTB) in the CNIC through a recording micropipette following electrophysiological characterization. Brains were sectioned and histochemically reacted to reveal both retrogradely labeled projection neurons in the DCN and anterogradely labeled descending projections from the IC. Contralateral reciprocal connections were examined between the labeled neurons via light and electron microscopy.

Results

Retrograde (CTB) and anterograde (BDA) label were readily identified at the injection site. CTB was found contralaterally in the DCN, whereas BDA was distributed bilaterally. Based on size and location, retrogradely labeled neurons comprise both pyramidal and giant cells. In the contralateral DCN, the close proximity of anterogradely labeled axonal swellings to retrogradely labeled cell bodies was highly suggestive of direct contact. Under electron microscopy, labeled axons were darkly stained and easily identified. Labeled soma were discernable due to accumulations of label within the Golgi apparatus and lysosomes. Synapses from descending projections were noted by the presence of a postsynaptic density on the retrogradely labeled cell adjacent to an anterogradely labeled terminal filled with synaptic vesicles. All labeled endings examined to date were filled with large round vesicles, suggesting the synapses are excitatory.

Conclusion

Our results demonstrate a direct tonotopic pathway by which the CNIC can bilaterally modulate activity in the DCN. Furthermore, our findings indicate that the contralateral descending projections are excitatory to the principal projecting neurons of the DCN. This point-to-point reciprocity provides a physical substrate for modulating ascending auditory information, and could be involved in “egocentric feedback” to enhance signal discrimination and/or selective attention.
Projections from the dorsal cochlear nucleus to the ipsilateral and contralateral inferior colliculi: single fiber reconstructions

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Background

Neurons of the dorsal cochlear nucleus (DCN) form a large projection to the contralateral inferior colliculus (IC) and a smaller projection to the ipsilateral IC. These projections are important because they may mediate sound localization based on spectral cues.

Methods

We labeled the projections to both the ipsilateral and contralateral IC by injecting the anterograde tracer, biotinylated dextran amine, in the DCN of mice. The trajectory, branching pattern, and synapses were studied.

Results

Injections resulted in many labeled fibers to the IC contralateral to the injected side and, in some cases, a few fibers to the ipsilateral IC. On the contralateral side, some fibers branched at the ventral edge of the IC, with a thicker branch innervating the central nucleus and dorsal cortex and a thinner branch innervating the lateral cortex. Some fibers only innervated the central nucleus and dorsal cortex. The few fibers that could be reconstructed to all their terminations demonstrated that a single fiber forms numerous branches with terminal and en passant swellings. On the ipsilateral side, the smaller number of labeled fibers (0-12 per case) made it easier to reconstruct fibers. Some of these fibers traveled mainly within the central nucleus along a presumed iso-frequency lamina. Other fibers formed terminations only in the lateral cortex. Tracing these ipsilateral fibers in the retrograde direction indicated that some of them were from bilaterally projecting axons. One terminal in the ipsilateral central nucleus formed an asymmetric synapse with round synaptic vesicles, characteristics of excitatory synaptic transmission.

Conclusion

Projections from the DCN to the contralateral and ipsilateral IC exhibited a variety of patterns, with some fibers innervating most of the IC and others targeting just one of its sub-regions. These diverse projections may indicate specialized patterns for specific aspects of sound localization as well as other functions. Support: NIDCD grant DC01089.
Perineuronal Nets on GABAergic Cells in the Inferior Colliculus that Project to the Medial Geniculate Body
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Background
Perineuronal nets are aggregates of extracellular matrix that can inhibit structural plasticity, promote synaptic plasticity, and protect neurons from oxidative stress (Karetko et al. 2009, Acta Neurobiol Exp 69:564). In many brain areas, perineuronal nets are preferentially associated with GABAergic neurons. In guinea pigs, nets in the inferior colliculus (IC) are most numerous in the central nucleus (ICc), though they also occur in dorsal cortex and external cortex. Most nets surround GABAergic cells, forming a population of “netted” GABAergic cells and a population of GABAergic cells without nets (Foster et al. 2012, Soc. Neurosci. Abst. 365.10). GABAergic IC cells project to a number of extrinsic targets; we test here whether nets are associated with GABAergic cells that project to the medial geniculate body (MG).

Methods
We injected red or green fluorescent beads or Fast Blue into the MG to label IC cells by retrograde transport. We stained GABAergic cells with antibodies to glutamic acid decarboxylase (GAD) and perineuronal nets with fluorescent-labeled Wisteria floribunda agglutinin. We identified retrogradely-labeled cells and determined whether they were GAD-immunopositive (i.e., GABAergic) and whether they were surrounded by a perineuronal net.

Results
A large injection in MG labeled GAD+ and GAD-negative cells in all IC subdivisions. Roughly 10-35% of the projecting GAD+ cells were surrounded by perineuronal nets. Small tracer injections demonstrated perineuronal nets surrounding IC GABAergic cells that project to each MG subdivision, with very few projecting to suprageniculate MG but a surprising number projecting to medial MG. While our focus was on GABAergic cells, we also observed IC-MG cells that were GAD-immunonegative (i.e., presumptive glutamatergic) and surrounded by perineuronal nets.

Conclusion
We conclude that perineuronal nets surround a subset of GABAergic IC-MG cells. These GABAergic projections terminate in multiple MG subdivisions. Nets also surround some glutamatergic tectothalamic cells. The proposed functional roles of perineuronal nets suggest that “netted” cells may react differently than other cells with regard to plasticity and oxidative stress. Future study of these topics in the IC may benefit from distinguishing not only GABAergic vs. non-GABAergic projecting cells, but also cells that have nets vs. those that do not. Supported by NIH DC04391 and DC012450.

Expression of the GABA\textsubscript{B}R1 and GABA\textsubscript{B}R2 Subunits in the Rat’s Inferior Colliculus
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Background
The type B \(\gamma\)-aminobutyric acid receptor (GABA\textsubscript{B} receptor) is an important GABAergic neurotransmitter receptor in the inferior colliculus (IC). GABA\textsubscript{B}R1 and GABA\textsubscript{B}R2 are two subunits of the GABA\textsubscript{B} receptor. One of our previous studies has revealed that the GABA\textsubscript{B}R2 subunit is expressed at a higher level in the dorsal cortex (ICd) than the external cortex and the central nucleus (ICx and ICc) of the IC. The present study was conducted in the rat’s IC to examine: 1) whether the GABA\textsubscript{B}R1 subunit had a similar area distribution; 2) how the overall level of the GABA\textsubscript{B} receptor in a specific IC region was dependent on the density of cell bodies expressing the receptor and the abundance of the receptor in the neuropil in the region; 3) morphological features of cells expressing the receptor.

Methods
Immunohistochemical experiments were conducted by using antibodies specific for the GABA\textsubscript{B}R1 and GABA\textsubscript{B}R2 subunits. Immunoreactivity was visualized using 3,3-Diaminobenzidine tetrahydrochloride reaction product and was quantified using optic density.

Results
GABA\textsubscript{B}R1 and GABA\textsubscript{B}R2 subunits were expressed in cell bodies and the neuropil throughout the IC. Regional distributions of the two subunits were similar to each other. For both subunits, the combined level of cell body and neuropil expression was higher in the ICd than the other subdivisions; and the area distribution of the expression in the neuropil was parallel to that of combined neuropil and cell body expression. The density of immunoreactive cell bodies tended to be higher but sizes of cell bodies tended to be smaller in the ICd than in the other subdivisions. No systematic regional
changes existed in the level of cell body labeling, except that GABA$_B$R2-immunoreactive cell bodies had slightly higher intensity of labeling in the ICd than in other regions. Elongated cell bodies existed throughout the IC. No strong tendency of orientation existed in immunoreactive cell bodies in ICc.

**Conclusion**

GABA$_B$R1 and GABA$_B$R2 subunits have similar area distributions in the IC, supporting that a functional GABA$_B$ receptor is a heterodimer. The contrast in the level of neuropil immunoreactivity among different subdivisions is consistent with the fact that the GABA$_B$ receptor has different pre- and postsynaptic functions in these different subdivisions. The lack of trend in the orientation of immunoreactive cell bodies in the ICc likely suggests that these cells are outside fibrodendritic lamiae in the structure, although this suggestion has yet to be verified by further experiments.

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**Non-Invasive Measures of the Health of the Implanted Cochlea**

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**Background**

The hypothesis that cochlear implant function depends on the health of the cochlea has motivated research and clinical efforts aimed at preservation and restoration of the normal cochlear biology. To help test the hypothesis we are examining several non-invasive, clinically applicable measures of implant function across animals that have a large range of cochlear pathology. Such measures are needed to assess the efficacy and stability over time of treatments designed to preserve or restore the health of the cochlea following implantation and to identify the healthiest stimulation sites when programming a patient’s processor MAP.

**Methods**

We define 'cochlear health' in terms of the percentage of surviving inner hair cells (IHCs), the density of spiral ganglion neurons (SGNs) in Rosenthal’s canal, and the level of ensemble spontaneous neural activity (ESA). We evaluated four potential monitors of cochlear health: electrically-evoked compound action potentials (ECAPs) and auditory brainstem responses (EABRs); and psychophysical temporal and multipulse integration functions. To create a variety of conditions in the implanted cochleae, four different treatment procedures were used in adult male guinea pigs. 1. Hearing group: implanted in a hearing ear, typically has retained IHCs. 2. AAV.NTF-3 treated group: deafened with neomycin, inoculated with an adenovirus containing an active neurotrophin gene to promote SGN survival and neurite regeneration. 3. AAV.Empty group: deafened with neomycin, inoculated with an empty AAV. 4. Deaf group: deafened with neomycin with no further procedures.

**Results**

ECAP data exhibited significant differences in slope steepness and level between animals with low versus moderate SGN densities. EABR data exhibited differentiation across all levels of SGN density. Both psychophysical monitors exhibited steep slopes and low thresholds only in animals that had >50% SGN density, moderate to good ESA levels, and surviving IHCs. Animals with no IHCs had shallow slopes (<1dB/doubling) and high thresholds, but some animals with surviving IHCs and ESA also had shallow slopes and high thresholds.

**Conclusion**

Cochlear health is reflected in the levels and slopes of the functions of these various non-invasive measures in guinea pigs. The psychophysical functions are good for differentiating between high and moderate to poor levels of cochlear health while the electrophysiological measures are sensitive to cochlear conditions across a large range of neural survival.
Listening Effort with Simulated Cochlear Implant Hearing and Electric Acoustic Stimulation – Effects of Noise on Response Time.

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Background
Cochlear implant (CI) fit is commonly assessed using speech intelligibility measures and self-report. However, these measures may not reflect listening effort. Prior research \cite{1} has shown that, when presenting normal-hearing listeners with CI-simulated speech, a dual-task paradigm can capture improvements in listening effort that are not reflected in speech intelligibility nor in subjective measures. These effects, however, were recorded near ceiling for speech intelligibility. The current study aimed to examine listening effort with simulated electric acoustic stimulation (EAS, combined CI and ipsilateral/contralateral acoustic hearing) compared to simulated CI alone, with intelligibility fixed at equal levels, below ceiling.

Methods
Normal-hearing listeners participated in one of three experiments. In experiment 1, intelligibility was fixed at 50\% sentence recognition (SRT50) and in experiment 2 at 79\% (SRT79) by presenting the speech stimuli at the appropriate signal-to-noise ratios (SNRs) in continuously present noise. In experiment 3 the stimuli were presented without noise, in alignment with the previous study \cite{1}. All three experiments used the same procedure, a dual-task paradigm proposed by Pals et al. \cite{1}. The participants listened to speech recordings processed to simulate CI hearing, residual low-frequency hearing, and EAS, and repeated back what was heard. Response times (RTs) on a simultaneous, visually-presented rhyme-judgment task served as a measure of listening effort.

Results
Preliminary results show no significant difference in RTs between simulated CI and EAS conditions presented at SRT50 or at SRT79. Average RTs for the CI and EAS conditions were about equal for both SRT50 and SRT79 and RTs recorded during listening to an auditory stimulus were shorter than RTs recorded between auditory stimuli. Preliminary results for experiment 3, without noise, show the reversed effect, in line with the previous study \cite{1}.

Conclusion
Noise affects the dual-task results in several ways; the difference in duration of RTs between and during listening is reversed compared to speech in silence, and the presence of noise may limit best performance on the response-time task. Other research has shown clear differences in listening effort between different intelligibility levels, yet the best RTs show no difference for SRT50 and SRT79. Preliminary results show no benefit in listening effort for simulated EAS compared to CI alone. However, effects of noise and listening effort need to be untangled before drawing final conclusions.

\cite{1} Pals, C., Sarampalis, A., Baskent, D., Listening Effort with Cochlear Implant Simulations, in revision.

Factors That Influence the Integration of Acoustic and Electric Stimulation in Bimodal Cochlear Implant Users

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Background
For cochlear implant (CI) users, effective combination of acoustic (A) and electric (E) stimulation may be limited by patient-related factors (e.g., residual hearing, performance in either ear), device-related factors (e.g., frequency overlap between devices), and/or listening condition (e.g., speech-in-noise, music). Likewise, optimization of combined electro-acoustic stimulation (EAS) may involve changing one or both device settings to address different concerns. This project aims to identify factors/conditions that limit the integration of electric and acoustic information for CI users’ speech and music perception.

Methods
Bimodal CI subjects, hearing aid (HA) in one ear and CI in the other, were tested using their clinical settings for both devices. Subjects were tested with the HA only (A), the CI only (E), and with both devices (EAS). Speech reception thresholds (SRTs) were adaptively measured for HINT sentences in speech babble. Melodic contour identification (MCI)
was measured with and without a competing instrument. The pitch of target contour and the competing masker was varied to test integration/segregation with E, A, and EAS.

**Results**

Preliminary results showed great inter-subject variability in terms of integration of EAS. A strong difference across speech and music was observed with EAS offering a greater benefit for speech, but little benefit or even interference for music. For most subjects, HINT SRTs were poorest with A-only, better with E-only, and best with EAS. Most subjects had difficulty with the MCI task with E-only (<50% correct), and most had no difficulty (>90%) with A-only at low frequencies. For music, some subjects attended to the better ear (EAS = A) while others had difficulty combining information across ears (EAS < A). For most subjects, performance was better when the pitch of the masker and target contours did not overlap; segregation using pitch cues was generally better with the A-only and EAS conditions than with E-only.

**Conclusion**

Acoustic hearing greatly allows for the perception of low frequency pitch cues which can be utilized in both speech and music. However, the present data suggest that the benefit of adding A to E depends on the listening condition, as well as patient-related factors such as the amount of aided hearing and the performance of the acoustic hearing ear. Such factors should be considered when optimizing the CIs and HAs for bimodal patients.

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**Influence of Insertion Site and Sealing Technique on Hearing Thresholds and Tissue Formation after Cochlear Implantation**

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**Background**

Nowadays severely hearing impaired people and patients suffering from profound hearing loss are treated with a cochlear implant (CI). Even though many of these patients benefit from receiving the implant, some challenges still remain. The development of a fibrous tissue sheath around the electrode carrier and/or new bone formation after implantation has an influence on the increase in impedance after surgery as well as on the hearing performance of the patients. Factors influencing the formation of connective tissue are the implant materials and surface properties as well as any trauma induced during surgery. As there is indication that the fastest and strongest growth of fibrous tissue and/or new bone formation is found always in the basal turn of the cochlea, the insertion site might have an influence on its formation. The aim of this study was to investigate the influence of the insertion site, round window membrane (RWM) vs. cochleostomy, and of different sealing techniques, no additional seal vs. muscle graft seal vs. bone cement seal.

**Methods**

Dunkin Hartely guinea pigs were implanted for 28 days with a silicone covered platinum-wire. All animals underwent frequency specific auditory brainstem response measurements (ABR) prior to surgery and at the end of the experiment to detect their hearing thresholds. Three groups were implanted through an incision of the RWM. The remaining groups received their implants via a drilled cochleostomy. Each implantation procedure was combined with three different sealing techniques (no additional seal, muscle graft seal, bone cement seal).

At the end of the experiment the cochleae were harvested and prepared for histology. Cochleae were embedded in epoxy and then a grinding procedure was performed (40 µm steps). After measuring the area filled with fibrous tissue/new bone formation, we calculated its volume. Additionally the ABR measurements were analysed to determine the threshold shift after implantation.

**Results**

In all animals a threshold shift after implantation was detected. In 1/3 of the animals implanted through the RWM the hearing threshold after surgery was above 90 dB SPL. In the groups implanted via a cochleostomy this occurred in 2/3 of the animals. Comparing the different sealing techniques, the groups with a bone cement sealing showed the largest threshold shifts as well as massive new bone formation.

**Conclusion**

According to our data, both, insertion site and sealing technique have an influence on the outcome after cochlear implantation.
Pre- and Post-Operative Binaural Masking Level Differences for Bimodal Cochlear-Implant Listeners

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Background

Cochlear implants (CIs) are increasingly recommended to individuals with bilateral acoustic hearing sensitivity that approaches or exceeds current clinical guidelines because of the documented benefits of combined electric and acoustic hearing. Although new electrode designs and surgical approaches are being developed to promote hearing preservation, most current patients will lose much of the acoustic hearing in the ear implanted. Consequently, there might be aspects of binaural hearing which they are sacrificing. While speech perception is generally improved from the combination of a CI with residual acoustic hearing, bimodal listeners have difficulty taking advantage of binaural squelch to improve speech understanding in noise. This study aimed to examine the potential trade-off between the benefits of a CI and the loss of binaural squelch for speech perception in noise as a result of cochlear implantation for individuals with limited bilateral acoustic hearing.

Methods

Binaural Masking Level Differences (BMLDs) were measured prior to unilateral cochlear implantation for 10 severely hearing-impaired subjects with some bilateral acoustic hearing (<80 dB HL at 500 Hz). Six of these subjects (to date) were also tested at least six months post-activation. IEEE sentences were presented audiovisually (including a video of the talker’s face) because auditory-alone performance was often too poor to reliably measure at signal-to-noise ratios (SNRs) where a BMLD would be expected. Speech signals were presented in speech-shaped noise for SNRs between −15 and +15 dB in 5–dB steps. The noise was always correlated between the two ears, while the speech signal was either correlated (N0S0) or inversely correlated (N0Sπ). Signals were delivered via headphones (95 dB SPL speech level) to the unaided ear(s) and/or via direct connection to the CI processor at MCL. BMLDs were calculated for each subject as the greatest dB difference between the N0S0 and N0Sπ psychometric functions.

Results

Four of 10 subjects showed evidence of a BMLD preoperatively (range: 2–8 dB). Of these four subjects, three have been tested postoperatively and none showed a measurable BMLD. However, all six subjects tested postoperatively showed N0S0 performance equal to or better than their preoperative N0Sπ performance.

Conclusion

Binaural sensitivity is diminished for some individuals with bilateral acoustic hearing who receive a CI, reducing binaural unmasking when the source of interest is spatially separated from the background noise. However, this deficit is likely overcome by the overall benefit to speech understanding provided by the CI.

Binaural Spectral Fusion in Hearing Aid and Cochlear Implant Users

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Background

Many hearing-impaired individuals receive little additional speech perception benefit or even interference from combining two hearing devices bilaterally, whether the devices are bilateral hearing aids (HAs), bilateral cochlear implants (CIs), or a CI worn with a HA in the contralateral ear (bimodal CI+HA). One contributing factor may be a change in the central processing of binaural inputs. Interaural pitch mismatches are often introduced by hearing loss, i.e. diplacusis, or CI programming, where real-world frequencies allocated to the electrodes differ from the electrically stimulated cochlear frequencies. Previous studies have shown that pitch perception adapts to reduce this discrepancy in some, but not all CI+HA users (Reiss et al., 2007). We hypothesize that some individuals instead adapt by increasing binaural fusion to reduce the perception of pitch discrepancy and, as in other sensory modalities, this fusion leads to averaging of information between ears.

Methods

Subjects were tested on three tasks: 1) Interaural pitch matching, in which a reference stimulus, a pure tone for a bilateral HA subject or a single electrode for a CI+HA subject, was pitch-matched to sequentially presented pure tones in the contralateral ear; 2) Dichotic fusion range measurement, in which a reference stimulus was presented simultaneously with a pure tone in the contralateral ear, and the contralateral frequency varied to find the frequency range that fused with
the reference;  3) Fusion pitch matching, in which a fused reference-contralateral stimulus pair was pitch-matched to sequentially presented tones in the contralateral ear. All stimuli were loudness balanced before testing.

Results
Seventeen CI+HA users and three bilateral HA users showed increased dichotic fusion frequency ranges of up to 2-3 octaves, compared to <0.03 octaves in normal-hearing listeners (van den Brink, 1976). The size of the fusion range was significantly correlated with the interaural pitch mismatch in both groups. In addition, averaging of dichotic pitch was observed, i.e. simultaneous presentation of two different pitches to the two ears resulted in the perception of a new binaural pitch intermediate between the original pitches.

Conclusion
The increased fusion frequency ranges and correlation of fusion frequency range with interaural pitch mismatch suggest that binaural fusion increases in some individuals as a compensation mechanism to reduce perceived interaural discrepancy. The increased fusion and associated spectral averaging between ears may account for speech perception interference effects observed with binaural compared to monaural hearing device use.

Residual Hearing Changes with Electrical Stimulation in Cochlear Implanted Guinea Pigs
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Background
Hybrid cochlear implants (CIs) are designed to preserve low-frequency (LF) hearing and allow combined acoustic-electric stimulation in the same ear. However, ~30% of these CI patients have >30 dB of LF hearing loss (HL) post-operatively between 3-36 months after CI activation (Gantz et al., 2010). The slow time course suggests that in addition to surgical trauma, electrical stimulation from the CI also contributes to HL. The goal of this study was to directly investigate the effects of electrical stimulation on residual hearing in a guinea pig CI model.

Methods
Twelve 2-month-old normal-hearing guinea pigs were implanted with an 8-ring animal electrode (Cochlear Limited, Australia) in the left cochlea via a cochleostomy. These animals were divided into two groups: chronic acoustic and electric stimulation (CAES; n=6) and no stimulation (NS; n=6). Non-implanted animals served as chronic acoustic stimulation controls (CAS; n=6). Auditory brainstem responses (ABRs) at 1, 2, 6, and 16 kHz and electrically-evoked ABRs (EABRs) were recorded biweekly to monitor changes in acoustic and electric hearing. In CAES animals, CIs were activated 5 weeks after surgery. EABR thresholds and behavioral responses were used to program T- and C-levels, respectively. CAES and CAS animals were stimulated 3 hours/day, 5 days/week for 10 weeks with modulated white noise. At the conclusion of the study, cochleae were processed and embedded in JB-4 resin for sectioning. Hair cell counts, stria vascularis vascular density, and spiral ganglion density were quantified.

Results
The majority of animals in the CAES and NS groups had significant post-surgical HL at 6 and 16 kHz. After 10 weeks of stimulation, four out of six animals in the CAES group had additional LF HL at 1 kHz. One out of six animals in the NS group had additional HL at 1 kHz; blood was observed in the apex of the cochlea for this animal. No additional HL was seen in the CAS group. Fibrosis, ossification, and changes in vascular density at the electrode region were seen in animals with post-surgical HL.

Conclusion
The greater incidence of increased HL at 1 kHz in the CAES group after electrical stimulation compared to the NS group suggests that electrical stimulation can contribute to HL with CI use over time. The results indicate that further study is needed to determine the effects of electrical stimulation on residual hearing.
REAL-TIME IMPEDANCE MEASUREMENT FEEDBACK FOR INTRACOCHLEAR ELECTRODE INSERTION

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Background
This pilot study details the use of a software tool that measures electrode impedance continuously during intracochlear electrode insertion, with the eventual potential to assess and optimize electrode placement and reduce insertional trauma.

Methods
A prototype software program for measuring electrode impedance and displaying it graphically in real time has been developed. The software was evaluated in both human cadaveric temporal bones and two live surgeries during intracochlear electrode insertion. Electrode positions were cross-evaluated in time alignment with the impedance measurements using real-time fluoroscopic analysis.

Results
Several electrode designs were evaluated with the new software system and data were obtained during electrode insertion with fluoroscopic guidance. Impedance changes were observed with variousicular positions. Using Contour Advance™ electrodes, impedance values increased after stylet removal, particularly when the electrodes are stimulated in the monopolar mode.

Conclusion
Impedance values seem to vary systematically with electrode position in scala; higher impedance values being associated with proximity to the cochlear wall. The new software is capable of acquiring impedance measurements during electrode insertion and this data may be useful to guide surgeons to achieve optimal and atraumatic electrode insertion, to guide robotic electrode insertion, and to provide insights about electrode position in the cochlea.

This study was supported by Cochlear Ltd. (PI: Roland). Dr. Tan’s participation in the study was supported by NIH grant K25-DC010834 (PI: Tan), and Dr. Svirsky’s participation was supported by NIH grant R01-DC003937 (PI: Svirsky).

Hyaluronic acid perfusion enhances cochlear implant insertion while preserving some degree of hearing function

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Background
Hyaluronic acid (Healon or HA) is frequently used as a substrate for the delivery of therapeutic drugs into the inner ear via the round window (RW) membrane. HA is also used as a lubricant in cochlear implant surgeries to facilitate a safer implantation. However, the effects of acute intra-cochlear exposure of HA on hearing are not well understood. HA is a highly viscous glycosaminoglycan polysaccharide that is widely present in human body and is bio-degradable. We have devised a novel cochlear implant insertion technique that entails perfusion of the cochlea with HA and takes advantage of its viscous nature. In present work we report (1) further developments in previously proposed implantation technique and (2) evaluate hearing function after perfusing the cochlea with HA.

Methods
A small cochleostomy was performed in the scala vestibuli (SV) of anesthetized, young adult Mongolian gerbils. RW membrane was removed to view the entrance to the tunnel of scala typani (ST). RW was filled with 1% HA and a mock-silicone implant was positioned directly above the entrance to the tunnel of ST with the help of a microcapillary. Another microcapillary was snugly inserted into the SV hole and a steady flow of HA was set up from RW to SV outlet. With the aid of the HA flow an implant was pulled into the cochlea. The effect of implantation and HA perfusion on hearing was assessed by comparing compound action potential (CAP) input-output functions and distortion-product otoacoustic emissions (DPOAEs) measured before and after the implant insertion. Perfusion with artificial perilymph using the same set up served as a control.

Results
Following artificial perilymph perfusion, CAP thresholds were comparable to pre-perfusion level. HA perfusion resulted in CAP threshold elevation that ranged from 30-60 dB across animals. However, none of the animals showed complete loss of hearing function following HA perfusion, which suggests that HA is not inherently ototoxic to hair cells and auditory-
nerve fibers. The CAP threshold elevations might have resulted from changes in mechanical function within the cochlea, which will be explored further. The new implantation technique resulted in the implant insertion all the way to the apex. Following HA perfusion, the DPOAEs were either reduced or in the noise floor with large variability across animals.

**Conclusion**
The new technique that flows an implant into the cochlea seems like a good alternative to traditional implant insertion methods.

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Long-Term Neurotrophic and Functional Effects of BDNF and Intracochlear Electrical Stimulation in Neonatally Deafened Animals

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**Background**
Postnatal development and survival of cochlear spiral ganglion (SG) neurons depend upon both neurotrophic support and neural activity. Our studies in cats deafened prior to hearing onset have shown that intracochlear electrical stimulation (ES) from a cochlear implant (CI) can help prevent SG degeneration, and intracochlear infusion of BDNF can further improve neural survival.

**Methods**
Kittens were deafened as neonates (systemic neomycin) and implanted at ≈1 month of age with custom-designed scala tympani electrodes with a drug-delivery cannula connected to a mini-osmotic pump. BDNF (94µg/ml; 0.25µl/hr) was infused for 10 weeks, and ES delivered for ≈5 months.

**Results**
Combined BNDF+ES elicited marked improvement in SG survival (>50% increase) and larger cell areas in the implanted ears as compared to contralateral. Moreover, SG survival after combined BNDF+ES was significantly better than after ES alone over these prolonged durations. BNDF+ES also resulted in angiogenesis in the SG, increased peri-implant fibrosis, higher density/larger size of radial nerve fibers within the osseous spiral lamina, and ectopic sprouting of these fibers into scala tympani. EABR thresholds improved relative to initial thresholds on chronically stimulated channels of the CI, but not on a control unstimulated channel.

In a separate study we asked whether neurotrophic effects would persist after cessation of BDNF infusion, **without ES**. After identical deafening and BDNF infusion procedures, animals were studied at ages similar to the BNDF+ES group (mean, 26 weeks). SG density data indicated that neurotrophic effects persisted for >3 months after terminating BDNF (38% increase). Substantial ectopic sprouting of radial nerve fibers again occurred.

Electrophysiological recordings in the inferior colliculus (IC) showed maintenance of the normal cochleotopic organization after BDNS+/-ES. IC thresholds were well-correlated with EABR thresholds. Lower IC thresholds were correlated with more selective central activation in all other groups, but not in BDNF-treated subjects where variability was much greater, perhaps due to the disorganized ectopic fiber sprouting.

**Conclusion**
BDNF promotes improved long-term survival of SG neurons and radial nerve fibers. These neurotrophic effects persist for ≥3 months after terminating BDNF delivery and withstand the additional stress of ES. Lower EABR thresholds and higher radial fiber density may be beneficial for CI function. However, the disorganized ectopic radial nerve fiber sprouting may be deleterious to optimized CI function.

Target structures for infrared neural stimulation (INS)
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Background
Pulsed mid-infrared lasers have been used as a method for neural stimulation, including cochlear stimulation. The use of lasers is appealing for many reasons, namely the spatial selectivity and non-contact method of stimulation. The mechanism of INS is believed to be involved with local heating of discrete neuron populations. In cochlear INS, some of the heat is absorbed in the cochlear fluid proximal to the stimulation site, resulting in measurable stress relaxation waves. These waves may vibrate the basilar membrane or directly stimulate hair cells, causing a stimulation by-product. Hair cell stimulation due to INS-created stress relaxation waves has been observed in vestibular system optical stimulation. The goal of the present study was to determine the contributions of hair cells to the recorded cochlear INS response.

Methods
In normal hearing animals, auditory responses can be masked with a noise stimulus. In the present experiments the INS was masked in hearing animals with an auditory stimulus. The optical fiber was inserted through a cochleostomy in the basal turn. After recording the effect of the masking using CAP measurements taken at the round window, the animals were deafened by 30mM Neomycin, which was injected directly in the scala tympani via the cochleostomy. After deafening was confirmed by testing for auditory-evoked CAPs, the masking experiment was repeated. If INS occurs through direct neural interaction, and not from hair cell stimulation, then the masking effect should disappear. Chronic deaf animals lacking hair cells were used in a second series of experiment, in which recordings from the central nucleus of the inferior colliculus (ICC) were used to determine whether stimulation of a cochlea lacking hair cells is possible.

Results
Preliminary data show that the masking effect is present in normal hearing animals and disappears after deafening with Neomycin. Animals that showed no hair cells in histologic sections revealed a neural response recorded from the ICC.

Conclusion
The effects of hair cells on INS can be seen through masking noise over optical stimulation. However, when hair cells are damaged or no longer present, INS still generates a neural response. These findings lead to the conclusion that although INS-induced stress relaxation waves may have some interaction with hair cells INS occurs through the direct interaction of the radiation and the neuron.

Optogenetic Stimulation of the Auditory Nerve: Toward an Optical Cochlear Prosthetic
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Background
When hearing fails, electrical stimulation of spiral ganglion neurons (SGNs) by cochlear implants (CIs) enables open speech comprehension in a majority of deaf subjects. Still, while healthy listeners encode a wide range of sound pressures (10^5) and frequencies (10^3 in 2400 steps), CIs typically employ only a narrow range of electrical currents to encode sound intensity and no more than 22 channels for frequency coding. The later limitation is caused by widespread current flow from each stimulating electrode, activating a large number of SGNs that represent many different sound frequencies. Optical SGN stimulation may improve CI coding by enabling more independent information channels to the CNS.

Methods
Here we employed channelrhodopsin-2 (ChR2), to render SGNs sensitive to blue light. We used transgenic mice expressing ChR2 under the Thy-1 promoter and virus-mediated expression of the ChR2 variant CatCh employing transuterine injection of AAV into the embryonic otocyst. Coupling blue light emitted by LED, into a cochleostomy or blue laser inserted through the round window, caused large compound potentials in scalp recordings.
Activation of the ascending auditory pathway was corroborated by light on tone masking and optogenetically evoked single neuron activity in the auditory nerve/cochlear nucleus and midbrain.

Results
Light stimulation caused compound potentials in scalp recordings (oABR) of hearing mice and in mouse models of acute and chronic human deafness. oABRs were absent in WT mice or in ChR2 mice when light was projected onto their intact cochlea. oABR generation was inhibited by lidocaine or TTX. Activation of the auditory pathway was corroborated by single neuron action potentials in the auditory nerve, cochlear nucleus and inferior colliculus. Light-on-tone masking corroborated specific activation of the ascending auditory pathway and coincided with the position of the cochleostomy relative to the tonotopic map of the cochlea as identified by high resolution X-ray tomography. Transuterine injections of CatCh-AAV into the otocyst effectively transduced SGN showing expression in SGN somata and neurites. oABRs and AP in the auditory nerve could be elicited in ears expressing CatCh.

Conclusion
- Optical stimulation of SGNs is feasible
- Specific activation of the auditory pathway was demonstrated by single neuron action potential from the auditory nerve/cochlear nucleus and inferior colliculus and by tone and light compound responses.
- AAV6-mediated expression of ChR2 constructs in murine SGNs is feasible.

Outlook
- Improve temporal bandwidth of stimulation by appropriate constructs.
- Study behavioral responses
- Develop mouse chronic intracochlear implants

Optogenetic control of central auditory neurons
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Background
Electric current spread limits the performance in users of central auditory prostheses by reducing the number of independent acoustic channels and increasing side effects from the activation of non-auditory pathways. Optogenetic stimulation of neuronal circuits has been proposed to improve the specificity of stimulation and has been demonstrated in a number of systems.

Methods
The left cochlear nucleus (CN) of CBA/J mice was transfected with AAV2/8-Channelrhodopsin-2 (ChR2) following posterior craniotomy in vivo. Following an incubation period of two to four weeks, mice were reanesthetized and blue light stimulation was delivered via an optical fiber placed near the surface of the transfected cochlear nucleus. Multi-channel recordings of neural activity were made in the inferior colliculus (IC) and in auditory cortex (Actx).

Results
A significant increase in neural firing rate was observed in the IC and Actx during blue light stimulation. Optical stimulation of the CN evoked spiking activity throughout the tonotopically organized central nucleus of the IC. This spread of activity was consistent with histological verification of transfection across the tonotopic gradient of CN. Varying pulse rate between 5 and 320Hz did not significantly affect threshold or bandwidth of the IC population response across animals. The spatial spread and magnitude of IC activity elicited by optical stimulation was comparable to that obtained with an acoustic click stimulus presented at 50 dB SPL. In contrast, no significant increase in neural firing rate was observed in response to optical stimulation in those cases with no ChR2 expression and in control cases. No acoustic artifact was measured when using the blue light laser.

Conclusion
These data suggest that optogenetic excitation of central auditory neurons is feasible and may provide the basis for a new generation of optically based neuroprosthetic devices.
Wavelength Dependency of Inner Ear Responses to Pulsed Laser Stimulation

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Background
Current research investigates the potential of optical stimulation of the inner ear to improve dynamic range and frequency resolution compared with stimulation via a conventional, electric cochlear implant. In addition to a potential for neural stimulation, laser radiation has been shown to cause pressure waves in the audible range along the axis of the laser beam in different media. In the current study we show cochlear sensitivity to nanosecond laser pulses at wavelengths ranging from ultraviolet to near infrared and for different pulse energies in vivo.

Methods
Cochlear responses to laser pulses were investigated by recording compound action potentials (CAPs) in seven normal hearing, anesthetized guinea pigs. The wavelength of the laser pulses were varied between 420 nm and 2150 nm for pulse energies between 3 µJ and 12 µJ. The stimulation site was located at the boundary between the basilar membrane and the spiral lamina seen through the round window. Both extra- and intracochlear stimulation approaches were investigated.

Results
We found a positive correlation between the energy level and the peak-to-peak CAP amplitude at all wavelengths investigated, with maximum CAP amplitudes comparable to ones evoked by acoustic stimuli at sound levels of 65 dB above threshold. These results could be shown for extra- and intra-cochlear stimulation. Stimulation between 420 nm and 2150 nm at constant energy levels led to a reproducible wavelength dependent CAP amplitude function. Below 1370 nm CAP amplitudes are significantly correlated to the absorption of either one of two main absorbers in the cochlea: hemoglobin and water respectively. Interestingly, the correlation with water absorption becomes negative for high absorption coefficients. We saw no CAP responses to laser irradiation following a Neomycin induced total hearing loss.

Conclusion
Overall laser evoked CAP characteristics are very similar to those of auditory evoked CAP. This is a good indication that laser stimulation can induce a substantial optoacoustic response in the cochlea. The strength of the response depends on pulse energy as well as the absorption coefficients of the media for the respective wavelengths. We see evidence for a change in the shape and extend of the absorbing volume at height absorption coefficients, leading to more localized optoacoustic effects and thus smaller CAP amplitudes than would be expected from the absorption strength alone. The extent of the observed optoacoustic effects demonstrates the importance of complete deafening in the investigation of neuronal laser stimulation.

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**Response Patterns to Infrared Neural Stimulation of the Cochlea Recorded from the Inferior Colliculus**

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**Background**

Cochlear spiral ganglion neurons can be stimulated with infrared radiation. While the stimulation is spatially selective and can be achieved over weeks without notable damages of the cochlea, it has to be demonstrated that infrared neural stimulation (INS) has the dynamics to encode the temporal structure of speech. In the present study, we further characterize the patterns of neuronal activity to irradiation of the cochlea with infrared pulses.

**Methods**

Single Tungsten electrodes and multichannel electrodes were placed in the central nucleus of the inferior colliculus (ICC) to recorded neural activity in response to INS of spiral ganglion neurons, both in normal hearing and deafened guinea pigs. Aculight tabletop diode infrared lasers (wavelength: 1844-1878nm) were used generate the optical pulses. The pulse length was between 10 and 200\(\mu\)s, the pulse repetition rate was between 1 and 250Hz, and the radiant energy was 0 to \(~126\mu J/pulse\). A 200\(\mu\)m optical fiber was inserted into the cochlea and was coupled to the laser to deliver the radiant energy to the target neurons. Optical stimuli included simple optical square pulses, triangular pulses, ramps up, ramps down, and amplitude modulated pulse sequences.

**Results**

Peri-stimulus histograms showed a single maximum with a latency of \(~5.4\) ms after the presentation of stimulus. Increasing the stimulus intensity shortened the latency of the first maximum to \(~5.1\) ms. Changing the stimulus pulse duration did not affect the response pattern. The maximum repetition rate for the neurons was lower for activity obtained with single Tungsten electrodes. For multi channel electrodes repetition rates of 200 pps could be seen.

**Conclusion**

The results for INS are similar to most of the data obtained with electrical stimulation. The data suggest that cochlear INS could provide stimulation with the appropriate temporal resolution to encode speech in future optical cochlear implants.

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**Modeling Performance of Optical Stimulation for use in Cochlear Implants**

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**Background**

Infrared neural stimulation (INS) has been demonstrated as an alternative to conventional electrical techniques for the stimulation of nerves. Based on the absorption of short pulses of infrared light, INS has the potential for higher spatial selectivity compared to standard electrical stimulation. There is interest in using INS to replace or complement electrical stimulation in bionic devices, specifically the cochlear implant where the increase in spatial selectivity would be advantageous. The majority of reports on INS have presented experimental results, whereas theoretical modeling of the processes involved can complement this work. Herein a detailed simulation of the localization of infrared stimulation, the potential for interference between multiple channels and the potential heat build up at high repetition rates is reported.

**Methods**

The model presented consists of a Monte Carlo model and finite element analysis (FEA) of the heat equation. Monte Carlo methods are utilized to simulate light transmission and absorption in tissue and provide spatial localization of an instantaneous laser pulse. The FEA solution of the heat equation reveals the temporal behavior of heat flow and allows repetitive pulses to be considered. The energy of the laser pulses and constants of the relevant media have been taken from the literature.

**Results**

The model predicts the temperature rise from single pulses can be as low as 0.01°C, depending on the wavelength and pulse duration, with the temperature decaying to the baseline temperature after \(~250\) ms. When a repetition rate of 10 Hz is used, the temperature peaks due to successive pulses overlap and the baseline temperature stabilizes at a value of
50% of an individual pulse's peak temperature. Faster repetitions result in greater temperature superpositions, a repetition rate of 100 Hz stabilizes at 760% of an individual pulse's peak temperature. When an array of spatially separated pulses are used, no increase in peak temperature is observed. However with a separation of less than 500 μm, the decay to baseline temperatures takes longer and an increase in the temperature superposition is observed over multiple pulse repetitions.

**Conclusion**

Results of INS modeling show that a significant increase in temperature is present in tissue when using a repetition rate similar to that used in current implants (> 100 Hz). The results show the need for further chronic experimentation to evaluate the effect of the increased temperature and show the potential advantage of a hybrid optical/electrical stimulation technique.

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**Water transport in the rat inner ear; translational research of Meniere's disease**

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**Background**

In order to develop new treatments of Meniere's disease, we investigated a time course of changes of the stria vascularis after vasopressin (VP) injection and the influence of OPC-31260 on experimentally induced enlargement of the intrastrial space caused by VP injection.

**Methods**

In the time course study, Wistar rats were injected with 50 mg/kg of VP subcutaneously. The stria vascularis specimens were harvested at 10, 20, 30, and 60 min after VP injection. For OPC-31260 administration, animals were administered 100 mg/kg of OPC-31260 orally 1 h before receiving 50 mg/kg of VP subcutaneously. The specimens were harvested 20 min after VP injection. These specimens were observed using transmission electron microscopy.

**Results**

In the time course study, the incidence of intrastrial space enlargement was 50%, 100%, 25%, and 0% for 10, 20, 30, and 60 min, respectively. In the study with OPC-31260 administration, the enlargement of the intrastrial space was reduced.

**Conclusion**

The intrastrial space was enlarged remarkably at 20 min after VP injection, and this enlargement of the intrastrial space was reduced by administration of OPC-31260 before VP injection. These results suggest that VP increases the influx of water from the perilymph to the basal cells via aquaporin (AQP) 2 and causes the formation of endolymphatic hydrops.

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**A Biodegradable Drug Delivery Microdevice for Tinnitus**

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**Background**

Tinnitus is the leading cause of disability in troops returning from Iraq and Afghanistan. It is often associated with blast injury and traumatic brain injury, and affects more than 13 million people in the US and Western Europe. Current treatment includes antidepressants and glutamate receptor antagonists. Here we report on a prototype biodegradable microdevice designed to deliver drugs to the cochlea for the treatment of tinnitus and other diseases of the inner ear. It is constructed from a recently developed class of biodegradable elastomeric poly(ester amide)s, poly(1,3-diamino-2-hydroxypropane-co-polyol sebacate)s (APS), showing a much longer and highly tunable in vivo degradation half-life compared with other commonly used biodegradable polymers.

**Methods**

A prototype implantable medical device fabricated from elastomeric scaffolds with tunable degradation properties is being developed for long-term drug delivery to the inner ear. Lidocaine is chosen as a surrogate for the early stage development
of the device, which will be designed to be compatible with emerging therapeutic compounds for the treatment of tinnitus. The microdevice is built by casting APS into porous Teflon molds, curing and assembling for the encapsulation of a therapeutic compound. The polymer degradation rate in vitro is measured compared to acceleration via enzyme degradation. Drug permeation properties of lidocaine through APS have been examined by performing Franz cell diffusion tests and simulated with MATLAB to predict release profiles.

**Results**

A prototype of the microdevice has been fabricated and tested in vitro. Lidocaine diffusion test shows that the diffusion coefficient of lidocaine through standard silicone is 10-fold higher than that of lidocaine through APS, allowing APS to better encapsulate lidocaine. Two different ratios of diamine:glycerol in the APS have been explored: 2:1 (diamine:glycerol) and 1:2, in order to establish optimal selectivity of the enzymatic triggering process relative to baseline degradation.

**Conclusion**

We report on a fully biodegradable drug delivery microdevice built using a fast, inexpensive, reproducible, and scalable fabrication process. The low diffusion coefficient of a small molecule surrogate (lidocaine), tunability in degradation rate, and selective degradation via lipase administration renders APS a suitable substrate for the delivery microdevice. When combined with emerging therapeutic compounds for tinnitus, this device will enable patient-controlled drug release via enzymatic degradation mechanisms at the desired time.

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### Pharmacokinetics of perilymph assessed with fluorescent markers

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**Background**

Understanding drug kinetics in the ear requires the use of quantitative procedures for drug delivery and fluid sampling and requires validated models for the interpretation of data. In the present study we have used sodium fluorescein and fluorescent dextran as markers to evaluate pharmacokinetic procedures.

**Methods**

Artificial perilymph containing 1 mM sodium fluorescein (NaF) or 1 - 2.5 mM fluorescein isothiocyanate conjugated dextran; FW 3000-5000 (F-dextran) was injected from pipettes sealed into the basal turn of scala tympani (ST), the cochlear apex, or the lateral semi-circular canal. As the cochlear aqueduct provides the outlet for injected volume, each procedure fills different regions of the ear. Either immediately after injection or after varying delay times, multiple fluid samples were taken sequentially from the cochlear apex or from the lateral canal. Each sample also represents perilymph from a different region of the ear. Perilymph sample concentrations were interpreted with the aid of a computer model.

**Results**

Previous studies have shown that NaF is a valuable marker as it is only slowly lost from perilymph. However, we found that following injection into the basal turn of ST, NaF was rapidly lost from ST to the middle ear through the round window membrane. Efflux was reduced when the round window membrane (RWM) was occluded with adhesive (Tissuemend) but was not blocked by inclusion of 5 mM probenecid, an organic anion transport inhibitor, in the injected solution. F-dextran applied into perilymph was consistently retained better than NaF but results with F-dextran applied into the basal turn of ST remain difficult to interpret. Perilymph concentrations were found to be dependent on the orientation of the animal, as a consequence of isosmotic F-dextran solution having a higher density than perilymph and “falling” along ST. Occlusion of the RWM also increased the retention of F-dextran.

**Conclusion**

Results with F-dextran show that pharmacokinetics of the basal turn of ST is more complex than previously appreciated. Distribution of F-dextran is affected by animal orientation, losses through the RWM, a sustained, slow entry of CSF via the cochlear aqueduct, distribution to adjacent compartments, distribution along the length of ST, and the elimination to blood. The data allow the parameters representing these kinetic processes in the computer model to be refined so that the model better represents the distribution of applied substances in perilymph of the inner ear.

This study supported by NIH/NIDCD R01 DC001368
**Inner Ear Gene Delivery with an EIAV-Based Lentiviral Vector**

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**Background**

Gene transfer into the cochlea provides a new opportunity for both research and future therapy for long-term protection, repair and regeneration. To date, a number of viral vectors have been evaluated for cochlear and vestibular gene delivery including adeno-associated virus (AAV), adenovirus (AV) and lentivirus. Lentiviral vectors have several advantages for cochlear gene therapy, including an ability to transduce terminally differentiated cells, long-term gene expression, large insert capacity (8-9 kb) and relatively early onset of gene expression following transduction. In this study, we evaluated the transduction efficiency of an equine infectious anaemia virus (EIAV)-based lentiviral vector carrying the green fluorescent protein (GFP) as a reporter gene (EIAV-GFP) in the inner ear of normal hearing mice.

**Methods**

We investigated several routes of administration including the posterior semicircular canal (PSC) via canalostomy, scala tympani (ST) through the round window and scala media (SM) via cochleostomy. Under general anesthesia, we administered EIAV-GFP (1µl) into the left ear via PSC, ST or SM. Two or 4 weeks after the surgery, cochlear and vestibular whole-mounts were stained for F-actin to identify the types of cells in the cochlea and examined by epifluorescence to determine the distribution of transgene expression.

**Results**

Two weeks after PSC administration, many cells in the non-sensory epithelium of the saccule expressed GFP, as did interdental cells in the cochlear basal turn. After ST administration, mesothelial cells showed GFP expression. SM administration resulted in GFP expression in several cochlear cell types, including fibrocytes in the osseous spiral lamina, pillar and Deiters’ cells, other non-sensory epithelial cells and the stria vascularis. Four weeks after administration, GFP expression was still present in all 3 treatment groups (PSC, ST and SM administration), although qualitative observation suggested the intensity of the green fluorescence was weaker than at 2-weeks.

**Conclusion**

EIAV-GFP transduces several different cell-types when administered via PSC, ST and SM inoculation. These data indicate that the EIAV-based lentiviral vector may be potentially useful for future clinical applications.

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**Using Gene Therapy Approach to Rescue Hearing in Connexin30 Null Mice by AAV1-Mediated Connexin Expressions in the Scala Media**

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**Background**

Non-sensory cells in the sensory epithelium of the cochlea are connected extensively by gap junctions (GJs) that facilitate intercellular ionic and biochemical coupling. Mutations in the genes coding for connexin 26 (Cx26) and Cx30 are the most common causes of human nonsyndromic hereditary deafness. Currently no mechanism-based therapy is available for treatment. We have previously identified a method to completely preserve hearing after directly inoculating viral vectors into the scala media when the surgeries were done in mice younger than postnatal day 5 (P5). In this study we performed gene-therapy studies using the Gjb6 null (Cx30 knockout) mice as the model.

**Methods**

GFP-tagged Cx30 (Cx30-GFP) or Cx30 without any tag was inserted into an adeno-associated virus (AAV) vector. These vectors were packaged into AAV2/1 viruses with the titers of 1.0x1013 and 1.5x1013, respectively. The recombinant viruses were injected into the scala media of one side of the cochlea of Cx30 null mice (P0 or P1). The contralateral side of the cochlea was not injected and used as a control.

**Results**

Without antibody labeling and beginning from 3 days after injections, we observed extensive green fluorescent punctas in the cell membrane of the claudius, outer sulcus and Hensen’s cells. Strong ectopic expressions in the cytoplasm of marginal cells, spindle-shaped cells were also observed. Weaker viral-mediated Cx30-GFP signal were detected in other
types of cochlear non-sensory cells as well. In Cx30 null mice expressed Cx30 without any tag, we observed similar pattern of expressions after labeling with the Cx30 antibody. Stronger expression was observed over time and the viral-mediated Cx30 expression lasted for at least 3 months, which was the longest time period we have observed so far. Exogenous Cx30 colocalized with endogenous Cx26. The normal function of these GJs was confirmed by fluorescence recovery after photobleaching assay. Dye transfer between GFP-positive cells showed more rapid fluorescence recovery than GFP-negative cells, although the rate of recovery was quantitatively slower than that measured from wild type mice.

Conclusion
Viral-mediated expressions of Cx30 with or without the GFP tag in Cx30 null mice were able to partially restore the intercellular communication mediated by gap junctions in the cochlea. Currently we are examining auditory brainstem responses to determine whether the viral-mediated expressions of Cx30 in the deaf Cx30 null mice were enough to rescue hearing.

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Development and System Integration of a Micro-actuator for Controlled Intracochlear Drug Delivery
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Background
Controlled, programmable intracochlear drug delivery may be necessary to treat certain hearing and vestibular disorders. Delivery directly to the inner ear using our system enables accurate dosing and sequencing of multiple compounds and overcomes obstacles such as the blood-cochlear barrier and the round window membrane. We have previously reported on our reciprocating system and in vivo testing for short and long-term experiments. In our design, a volume of solution (perilymph and concentrated drug) is repeatedly infused and then withdrawn, resulting in zero net volume transfer to the scala tympani while allowing drug transport by diffusion and mixing within the perilymph. Further miniaturization and functionality is required for an implantable device targeted for human therapeutic use.

Methods
A custom micro-actuator drives a reciprocating pump and reservoir refresh pump. It has been developed to replace the larger, heavier commercial actuator and pump currently in the system. We are prototyping low-power electromagnetic assemblies which are surface-mounted to our planar, laminated fluidic manifolds. The assemblies interface to membranes which are part of either pumping chambers or valve structures. The actuators are electronically controlled to provide variable pumping rates and valve sequence control. We are also integrating fluidic check valves as part of our pump development effort. Parallel development of the reciprocating pump and reservoir refresh system proceeds toward long-term bench testing and sustained in vivo drug delivery.

Results
The actuator (with a volume of less than 0.2 cc) is sized to fit the form-factor constraints of the human implantable device. It is designed with dual electromagnetic and permanent magnetic control to achieve low-power operation. It can produce up to 1 N of force over a displacement of several hundred microns, generating fluid displacements between 0.1 and 1.0 microliter per cycle. The actuator requires very little power, using approximately 1 mJ of energy or less per actuation cycle. Additionally, the actuator is utilized, in combination with in-line valves, to drive the positive-displacement reservoir pump. Integration of the actuator with fluidic components and test results at the system level have been obtained.

Conclusion
Low-power, miniature electromagnetic actuators will be required for reciprocation and reservoir refresh in order to meet the size requirements of clinical systems. These results represent an advance in miniaturization and enhanced functionality towards the goal of a fully implantable drug delivery system for treatment of SensoriNeural Hearing Loss and other diseases.
Intracochlear Tamoxifen Distribution Visualized by a Cre-Lox System
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Background
Local delivery to the cochlea has the potential to avoid systemic side effects of drugs such as steroids, and has resulted in widespread use of transtympanic approaches to drug delivery in the clinic. Detailed pharmacokinetic studies of drugs applied to the inner ear have relied on the determination of drug concentration in perilymph, which is technically challenging in animal models and does not reveal whether drug gains access to individual tissues or cells. We have developed a novel approach to the determination of drug distribution at the cellular level using a Cre-Lox system in which we can visualize exposure of individual cells to tamoxifen, a synthetic steroid analogue.

Methods
We used two Cre-reporter strains to perform cell tracing. To assess exposure to cochlear supporting cells we used a Cre strain driven by the Sox2 promoter and for visualization of all exposed cells we used a Cre strain driven by the CMV promoter, which is ubiquitously expressed and labeled the cochlear lateral wall, spiral limbus, and spiral ganglion cells. The Cre strains contained fusion proteins with the estrogen receptor rendering them sensitive to tamoxifen. The mice were crossed to a reporter strain that converts from expression of tdTomato, a red fluorescent protein, to expression of GFP upon Cre activity. We used several doses of intraperitoneal tamoxifen as the systemic treatment, and delivery of drug via the round window niche or lateral semicircular canal for local delivery. The number of GFP-positive cells was counted in ears dissected 1 week after drug treatment.

Results
A clear dose-dependent response to systemic tamoxifen injection was observed in all cochlear tissues. Tamoxifen showed a gradient of distribution from the basal to apical turns of the cochlea after delivery by the round window niche approach. No significant differences in the location of GFP-positive cells along the cochlear spiral were observed by the posterior semicircular canal approach.

Conclusion
We could visualize tamoxifen distribution in cochlear tissue with the use of a novel reporter system. Since cells retained GFP expression after exposure to tamoxifen, timing and distribution of tamoxifen could be monitored in the cochlear tissue. This reporter system could be a valuable strategy for the visualization of drug distribution in any organ.

Microparticle Characterization for Cochlear Drug Delivery
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Background
Cochlear implants are the treatment of choice for patients with moderate to profound sensorineural hearing loss, however increasing numbers of cochlear implant recipients have remaining hearing that needs protection from insertion trauma. Local delivery of therapeutic agents could potentially protect remaining sensory cells and neuronal connections. The ideal carrier for local delivery would have the ability to be functionalized to attach to the cochlear implant and/or cochlear cells, deliver multiple drugs with distinct pharmacokinetics, and sustain factor release over a defined period of time. It would also be useful to be able to monitor carrier distribution in the cochlea via fluorescent probes incorporated into/onto the carrier in test studies. To that end, fluorescent multicompartimental microparticles appear to be excellent candidates for use as carriers for drugs aimed at reducing cochlear implant trauma. Specifically, we characterize the feasibility of intrascalar delivery, detection of multicompartimental microparticles in the cochlea, and the controlled drug release from the particles.

Methods
Particles were fabricated from an FDA approved biodegradable polymer and a polysaccharide via a process called electrohydrodynamic co-jetting. For in vivo assessment, 5µL of 15mg/mL microcosmic solutions were infused via cochleostomy into the scala tympani of a guinea pig animal model at a rate of 1µL/min. For in vitro experiments, Piribedil, an anti-excitotoxic drug, was incorporated into the polymer matrix at 2% w/w (drug to polymer in one compartment) prior to jetting. Confocal laser scanning microscopy (CLSM) was used to confirm particle composition and tricompartmental nature, scanning electron microscopy to determine particle size, shape and surface topology, and 2 photon CLSM to delineate in vivo particle distribution within the 3D volume of the cochlea. In vitro drug release was assessed with UV
spectrophotometry. After in vivo infusion, cochleae were harvested, decalcified, and sectioned to enable visualization of particles within the scala tympani and cochlear tissue via CLSM.

Results
Microparticles composed of distinct compartments were present in the cochlea up to 24 hrs post-infusion. The majority of the particles were found in the basal turn near the site of the cochleostomy. Piribedil was successfully incorporated into the microparticles and the release profile over a period of 14 days was determined.

Conclusion
Multicompartmental microparticles have demonstrated promise as intracochlear drug carriers as they exhibit controlled drug release and the ability to persist in the cochlea.

Sustained release glucocorticoid formulation for long term intracochlear drug delivery
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Background
Previously we reported our short term data on extended release glucocorticoid (GC) formulations using various dose-ranges. Our work demonstrated very good short term sustained release profiles in vitro and in vivo and studies in animal models demonstrated excellent safety. One major advantage of the O-Ray drug delivery system is the ability to deliver consistent amounts of therapeutics over very long periods of time.

Methods
To show this in vivo, O-Ray extended release GC formulations (predicted 100 day release) were implanted into guinea pig cochleae. Pre- and post- treatment otoacoustic emissions and ABRs were performed and excellent hearing preservation was observed. Pharmacokinetic studies were performed testing GC presence in the perilymph at 90, 120, and 180 days.

Results
Interestingly, superior hearing was observed in ears administered extended release steroids at four and six months after implantation, possibly suggesting a rescue of age dependent hearing loss. As expected, all ears containing an implant showed significant GC levels in the perilymph at 90 days (5/5). Some animals still had detectable GC in the perilymph at 120 days (2/4) and 180 days (1/4) showing the potential for even longer sustained release durations.

Conclusion
In otology, GCs are routinely used to treat sudden hearing loss, Meniere's disease, autoimmune inner ear diseases, and certain vestibular disorders. Despite their widespread use, GCs can cause serious systemic side effects and a localized sustained release formulation of GCs is expected to be more effective and safer for use in the ear. The ultimate goal of this study is to develop a tunable system for the delivery of glucocorticoids to the ear for the treatment of myriad diseases of the inner ear.
Rescue of cisplatin dependent hearing loss by an intracochlear drug delivery implant
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Background
Chemotherapy induced ototoxicity is damage to the inner ear caused by side effects of administered drugs. This can lead to permanent hearing loss and/or tinnitus in the patient. Some of the most common drug classes causing ototoxicity are anti-cancer therapeutics such as cisplatin. Cisplatin is considered one of the most ototoxic drugs in use, with monitored ototoxicity frequencies commonly ranging from 20-100\% depending on the dosage used. The development of an otoprotective treatment to combat chemotherapy induced hearing loss would have significant impacts on patient quality of life. Various compounds have been studied and tested in animal models for protection against cisplatin mediated hearing loss. Among those with some ototoxic activities include antioxidants, sulfur containing nucleophiles, and steroids. Steroids remain attractive candidates as they are well characterized, and have been used extensively for various inner ear indications.

Methods
We previously reported our extended release steroid formulations in various dose-ranges utilizing a sustained release polymer formulation. To test their activity, sustained release steroid formulations were implanted directly into the cochleae of guinea pigs followed by a cisplatin challenge sufficient to induce hearing loss.

Results
Pre- and post- treatment otoacoustic emissions and ABRs were performed to assess hearing. In all cases, significant hearing improvement was observed in ears administered steroid implants.

Conclusion
These results demonstrate otoprotection from cisplatin ototoxicity and form the foundation for the initiation of clinical trials.
Administration of Glucocorticoid Attenuates Trauma-induced Hearing Loss Associated with Intracochlear Cell Transplantation

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Background
One of the greatest challenges in the treatment of inner ear diseases is to find a cure for hearing loss caused by the loss of cochlear hair cells or spiral ganglion neurons. Development of stem cell transplantation raises the hope for future regeneration therapy in the inner ear. However, residual hearing usually deteriorates after intracochlear cell transplantation. Application of glucocorticoid has been shown to ameliorate hearing loss resulting from the trauma of cochlear implantation in experimental animals. The purpose of this study is to investigate whether administration of glucocorticoid can protect inner ear from hearing loss resulting from the trauma associated with intracochlear cell transplantation.

Methods
Embryonic stem cells were delivered into the cochlear via the round window membrane of B6 mice. Betamethasone of either 200 µg/ml or 800 µg/ml was injected into the cochlea with the transplanted stem cells simultaneously. Auditory brainstem responses (ABRs) were measured immediately, 1 week, and 2 weeks after the cell transplantation. After 2 weeks, the distribution and survival of the embryonic stem cells were investigated by immunohistochemistry study. Terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) assay was used to detect the possible apoptotic event in the treated inner ears.

Results
The ABR thresholds increased immediately after the cell transplantation in both the control group and the betamethasone-treated groups, and the hearing partially recovered in the treated groups at 1 week and 2 weeks after the surgery. Simultaneous injection of 800 µg/ml betametasone with the transplanted significantly attenuated the hearing loss resulting from the trauma associated with intracochlear cell transplantation.

Conclusion
Simultaneous administration of glucocorticoid diminishes the trauma caused by intracochlear cell transplantation. These findings might provide insight into a novel strategy for preserving the residual hearing when stem cell therapy is to be applied via the intracochlear route.

Identification of COCH gene mutation in exon 5 of the LCCL Domain in Archived DFNA9 Temporal Bone

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Background
DFNA9 is an autosomal dominant nonsyndromic hereditary hearing loss due to heterozygous mutation in the COCH gene, encoding cochlin. Manifestations of DFNA9 include adult onset progressive sensorineural hearing loss and vestibular dysfunction. Cochlin is the most abundant protein in the inner ear. Our laboratory and others have previously described the histopathology of DFNA9 at the otopathological level. Interestingly, the late manifestations of DFNA9 include cartilage deposits in the tympanic membrane, mucosa, and ossicular joints. However, anacusis precedes the dysfunction of the ossicular chain or tympanic mobility, and patients do not typically present with a mixed hearing loss.

Methods
Extraction of genomic DNA from archived celloidin-embedded temporal bone was performed. This temporal bone was previously identified as having an LCCL domain mutation via proteomic analysis in our laboratory. Temporal bone DNA and control DNA (commercial human genomic DNA) were amplified using PCR with primers designed to flank exon 5 of
the LCCL domain, spanning the region corresponding to the suspected mutation. PCR products of 396 bp amplicon length were gel purified and sequenced.

Results
Sequencing of PCR products from the DFNA9 temporal bone revealed a heterozygous single nucleotide G->A transition at residue 6328 within exon 5, which was predicted to result in an A119T (Ala to Thr) amino acid change in the cochlin protein product.

Conclusion
We have verified a mutation in the COCH gene that we previously identified at the protein level via proteomic analysis from an archived formalin fixed celloidin-embedded temporal bone. At the time this temporal bone was received, our laboratory was not routinely collecting another DNA source from donors; whereas, we now collect buccal swab samples. To our knowledge, this is the first report of an hereditary hearing loss mutation identified in archived temporal bone tissue.

Diagnostic application of targeted exome sequencing for familial nonsyndromic hearing loss
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Background
Identification of causative genes for hereditary nonsyndromic hearing loss (NSHL) is important to decide treatment modalities and to counsel the patients. Because the genetic heterogeneity is extraordinarily huge in sensorineural disorders, we need to develop high-throughput diagnostic tools. To this end, we designed molecular genetic diagnosis based on targeted exome sequencing (TES) to screen all the candidate genes for NSHL.

Methods
Targeted exome capture of 80 known deafness genes and massively paralleled sequencing were done in 20 probands from families with two or more members with variable degree of hearing loss. To identify the causative mutations among variants found in TES, we filtered out the variant that were not compatible with the inheritance pattern of the family. Finally we validated all the candidate variants by Sanger sequencing. We further checked the variants if they are detected among normal hearing 80 control subjects. Genotype-phenotype correlation including configurations of audiogram and the progression pattern of hearing loss was also used.

Results
Critical mutations were identified in 11 of 20 probands of these multiplex families through NSHL80-TES. We identified damaging mutations in nine genes such as WFS1, COCH, EYA4, MYO6, GJB3, MYO6, COL11A2, MYO3A and MYO7A. Most of them are private, again confirming etiologic heterogeneity of hereditary deafness in this population. In this study, we suggested the NGS-based new diagnostic flow to identify mutations responsible for familial NSHL, which detected mutations in 55% of multiplex families with hearing loss.

Conclusion
We suggested a diagnostic flow to identify a causative mutation responsible for hearing loss. This genomic analysis procedure will enable us to evaluate causative mutation with more efficiency.
Differences in the Pathogenicity of the p.H723R Mutation of the Common Deafness-Associated SLC26A4 Gene in Humans and Mice

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Background
Recessive SLC26A4 mutations contribute to both Pendred syndrome (PS) and nonsyndromic hearing loss (DFNB4), constituting a common cause of hereditary hearing impairment worldwide. In recent years, the understanding of the pathogenesis of PS and DFNB4 has been accelerated by various mouse models with mutations in the Slc26a4 gene, including knock-out Slc26a4-/- mice, Slc26a4loop/loop mice with the p.S408F mutation, Slc26a4tm1Dontuh/tm1Dontuh mice with the c.919-A>G mutation, and conditional knock-out Tg[E];Tg[R];Slc26a4£G/£G mice. However, none of these mouse models could perfectly simulate the progressive or fluctuating hearing loss in humans. In the present study, we established a knock-in mouse model with the p.H723R (c.2168A>G) mutation, a common SLC26A4 mutation in the Asian population, and then, we characterized the associated audiovestibular phenotypes as well as the inner ear pathology.

Methods
Mice homozygous for the p.H723R mutation (i.e., Slc26a4tm2Dontuh/tm2Dontuh) were generated and maintained in the C57BL/6 background. Corresponding to the human genotypes, mice with compound heterozygous mutations for p.H723R and c.919-2A>G (i.e., Slc26a4tm1Dontuh/tm2Dontuh) were also obtained. Mice were then subjected to audiological assessments, a battery of vestibular evaluations, inner ear morphological studies, and noise exposure experiments. The expression of pendrin was examined using immunolocalization; whereas the expression of Kcnj10, a gene regulated by Slc26a4 and involved in the pathogenesis of Slc26a4 mutations, was investigated using real-time PCR and Western blotting.

Results
Both Slc26a4tm2Dontuh/tm2Dontuh and Slc26a4tm1Dontuh/tm2Dontuh mice showed normal audiovestibular phenotypes and inner ear morphology, and they did not show significantly higher shifts in hearing thresholds after noise exposure than the wild-type mice. The expression of pendrin and Kcnj10 was not affected in Slc26a4tm2Dontuh/tm2Dontuh and Slc26a4tm1Dontuh/tm2Dontuh mice as compared to the wild-type mice. The results indicated that the p.H723R allele was non-pathogenic in mice because a single p.H723R allele was sufficient to maintain normal inner ear physiology.

Conclusion
Using a genotype-driven approach, we generated a knock-in mouse model segregating the common deafness-associated SLC26A4 p.H723R mutation in humans. To our surprise, mice with the Slc26a4 p.H723R mutation had a normal audiovestibular phenotype and inner ear morphology. Because there might be differences in the pathogenicity of specific SLC26A4 mutations in humans and mice, precaution should be taken when extrapolating the results of animal studies to humans.
Large-scale Application of Next Generation Sequencing Approach in Screening Mutations of Currently-known and Candidate Human Deafness Genes in Multiple Ethnic Populations

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Background
Targeted capture and sequencing of all reported human genes linked to deafness by the next-generation sequencing (NGS) method may provide a powerful tool for studying the spectrum of genetic mutations in different human ethnic populations. The uses of the NGS technology for mutation screening of deafness genes on an epidemiological scale have been hindered by (1) a bottleneck in streamlined samples processing; (2) a high per-sample cost; (3) difficulties in bioinformatic data analyses.

Methods
In order to solve these three major problems we have developed: (1) a flexible robotic process for NGS library preparation based on the Hamilton robotic liquid handling system; (2) a low-cost hybridization-based gene capture method for enriching exons of >120 genes that have been reported to cause deafness in humans and mice; (3) a bioinformatic pipeline for automatic identification of disease-causing mutations from large numbers of samples and generation of reports.

Results
The automatic library preparation has the flexibility of processing 1 to 96 samples in one run, and generated quality of Illumina libraries similar to that done manually. Our hybridization-based gene capture method achieved specificity, multiplexicity, uniformity and depth of coverage in exon captures suitable for downstream Illumina sequencing. The high coverage depth and cost benefits of our approach is applied for genetic screening of close to one thousand samples so far, including samples collected from Asia (Han and Tibetan Chinese, Korean), Africa (Ivory Coast and Cameroon) and the United States. Samples from confirmed cases of genetic deafness, from unknown etiology but with genetic suspicion and from clinically-confirmed normal hearing subjects were tested. Preliminary data obtained found a spectrum of mutations in GJB2, SLC26A4, WFS1, DFNB59, GJB4, CDH23, KCNE1, MYO15A, KCNJ10, MYO1F, GJB3, and POU3F4 genes. Further bioinformatic data analysis is ongoing.

Conclusion
We are approaching the milestone of developing an automatic pipeline for the large-scale application of NGS for genetic screening of about 120 deafness genes for clinical applications. We will report our findings from these multi-ethnic populations in terms of the rank of most prevalent genetic mutations found, the error rates in terms of both false negatives and false positives, and confirmation studies with Sanger sequencing.
Loss of FGF23 signaling results in mixed hearing loss and middle ear malformation

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Background

Fibroblast Growth Factor 23 (FGF23), a member of the FGF19 subfamily, is a circulating hormone, secreted by osteoblasts and osteocytes, which regulates renal phosphate handling. KLOTHO is a single-pass transmembrane protein that acts as a critical cofactor for FGF23 mediated signaling by increasing FGF Receptor (FGFR) affinity for FGF23, and is believed to account for FGF23’s tissue specific function. KLOTHO expression has been observed in the cochlea, specifically in the stria vascularis, spiral ligament, OHCs, IHCs and SG cells. KLOTHO−/− mice exhibit hearing loss, and are used as a model of age-related hearing loss. Cochlear histology in KLOTHO−/− mice demonstrates no obvious morphological abnormalities and normal SG neuron densities.

Multiple hypotheses exist concerning the mechanism of hearing loss in KLOTHO−/− mice; it is unclear whether or not the KLOTHO−/− phenotype is independent of FGF2 signaling. Motivated by the critical role that several FGFs play in the developing auditory organ, and the potential implications of the KLOTHO−/− mouse, this work sought to uncover the role FGF23 plays in the auditory system.

Methods

The physiological performance of the auditory system in FGF23−/− and FGF23+/− mice was assessed using ABR and DPOAE measurements. Anatomical differences were assessed via histological and morphological characterization of the bulla, ossicles and cochlea using light microscopy and high-voltage X-ray computed tomography (micro CT).

Results

ABR measurements from FGF23−/− mice demonstrated profound hearing loss across the physiological frequency range, while FGF23+/− mice had statistically normal hearing below 20 kHz, and losses of up to 25 dB above 20 kHz. Similar to ABR results, DPOAE assessment of FGF23−/− mice demonstrated nearly complete hearing loss, while FGF23+/− mice had severe loss at the highest frequencies, and nearly normal hearing below 20 kHz.

Morphological assessment of middle ears in FGF23−/− mice demonstrated cartilaginous bone and malformation of the bulla and ossicles; FGF23+/− mice had near-normal morphology. The cochleae and vestibular organs of FGF23−/− and FGF23+/− mice appeared normal grossly, and on microscopic analysis of midmodiolar sections.

Conclusion

FGF23−/− mice have profound hearing loss, and extensive middle ear malformations. FGF23+/− mice have mixed hearing loss. The observed hearing phenotype does not match observations in KLOTHO−/− mice, and suggests that loss of FGF23 signaling impacts the cochlea via different mechanisms than KLOTHO.
Interaction between Adiponectin and Adiponectin Receptor 1 is Associated with Age-related Hearing Impairment

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Background
Age-related hearing impairment (ARHI) is a complex disease caused by an interaction between environmental and genetic factors. Recently, several studies confirmed obesity as an independent risk factor for ARHI. In our previous study investigating the underlying mechanisms, we demonstrated that plasma adiponectin might protect peripheral hearing function. It has been revealed that polymorphisms of the adiponectin gene, ADIPOQ, might affect plasma adiponectin levels; and polymorphisms of both ADIPOQ and its type 1 receptor gene, ADIPOR1, have been related to obesity-related morbidities. Hence, we postulated that genotypes of ADIPOQ and ADIPOR1 might be associated with the development of ARHI.

Methods
A total of 1682 volunteers (Han Chinese, aged 40 to 80 y) were included in the clinical analyses, and their audiological phenotypes were determined according to the Z scores converted from their original frequency-specific hearing thresholds. By using the database of Chinese Han haplotypes in International HapMap Project and NIEHS, followed by analyses with the haploview software, 9 tagSNPs and 4 tagSNPs in ADIPOQ and ADIPOR1, respectively, were selected for genotyping. The genotypes were then correlated to the audiological phenotypes under the assumption of various inheritance models.

Results
The rs2241767 [G] allele and the rs1063539 [C] allele of ADIPOQ were significantly associated with low-tone Z scores under dominant, additive, and co-dominant models; whereas no association was identified between the ADIPOR1 variant alleles and Z scores. To explore the interaction between ADIPOQ and ADIPOR1 concerning the contribution to ARHI, we stratified the subjects according to their genotypes of the ADIPOR1 tagSNPs, and then correlated ADIPOQ genotypes with the Z scores in each strafication of subjects. Of note, the association between rs2241767 and rs1063539 of ADIPOQ and Z scores appeared to exist in subjects with certain ADIPOR1 genotypes only. These findings were then validated on haplotype analyses.

Conclusion
ADIPOQ variant alleles were associated with ARHI. In contrast, although ADIPOR1 variant alleles were not directly associated with Z scores, the association between ADIPOQ and hearing thresholds seemed to be modulated by the ADIPOR1 genotypes. In other words, the development of ARHI might result from an interaction between adiponectin and type 1 adiponectin receptor.
Novel pathological model of SYM-1 and conductive hearing loss revealed by docking simulation of Noggin and heparin
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Background
The access of bone morphogenetic protein (BMP) to the BMP receptors on the cell surface is regulated by its antagonist Noggin. Noggin-BMP complex associates with heparan-sulfate proteoglycans (HSPGs), a major proteoglycan on the cell surface and in extracellular matrices, and is regulated by sulfatases to control the local activity of BMPs. Mutations in NOG are associated with various autosomal dominant syndromes that are characterized by a spectrum of skeletal defects and synostoses, such as multiple synostoses syndrome (SYNS1), Teunissen-Cremers syndrome (TCS), and tarsal-carpal coalition syndrome (TCC). Proximal symphalangism (SYM1), SYNS1, and TCS are usually associated with conductive hearing loss. It is not yet understood how different NOG mutations can produce this wide range of symptoms.

Methods
Genetic analysis of NOG gene of a family with SYM1 and conductive hearing loss was performed. To compare normal and R136C mutant Noggin structure, we generated modeling structures using crystal structure as a template. To analyze the effect of R136C on the function of Noggin, we conducted docking simulation of the Noggin and pentasaccharide analogue of heparin (used as the ligand) using AutoDock4.2.

Results
A pedigree diagnosed as SYM-1 was found to have a novel heterozygous missense mutation (p.R136C) associated with the symptoms. The position 136 was the heparin-binding site of Noggin, raising the possibility that R136C would change the binding affinity. No mutations of the heparin-binding site of Noggin have previously been reported to associate with diseases. We utilized the crystal structure of wild-type Noggin and investigated whether the p.R136C mutation altered its structure that could lead to some pathogenic effect. An in silico docking analysis showed that the heparin analogue appeared to associate with the putative heparin-binding site (K133-K144) of wild-type Noggin. However, one of the salt bridges between Noggin and heparin pentasaccharide disappeared following the replacement of the arginine at position 136 with a non-charged cysteine.

Conclusion
Based on our computational analyses, we predicted that the number of mutant Noggin-BMP complexes that are tethered to HSPGs on the cell surface are decreased because of the low binding affinity of the Noggin with R136C mutation to HSPGs. Reduced binding ability of the mutant Noggin to HSPGs is likely to the cause of excess BMP signaling that ultimately resulted in PIP joint fusion and conductive hearing loss.
of Mcph1 in the epithelial cells of middle ear cavities supported its involvement in the development of otitis media. Other defects of Mcph1tm1a/tm1a mice included small skull sizes, increased micronuclei in red blood cells, increased B cells and ocular abnormalities.

**Conclusion**

These findings not only recapitulated the defects found in other Mcph1-deficient mice or MCPH1 patients, but also revealed an unexpected phenotype, otitis media with hearing impairment, which suggests Mcph1 is a new gene underlying genetic predisposition to otitis media.

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**Targeted Exome Capture and Paired-End Massively Parallel Sequencing Reveals New Mutations for Human Hereditary Deafness in the Middle East**

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**Background**

Classic techniques such as linkage analysis and Sanger sequencing have led to the discovery of over 100 genes for deafness, a highly genetically heterogeneous disease. Nevertheless, a significant portion of hereditary hearing loss remain unsolved and many more genes remain to be discovered. This is particularly true of the Middle Eastern population, with many different ethnic groups and high rates of consanguinity. Exome capture and massively parallel sequencing can be exploited to address this challenge.

**Methods**

A targeted capture pool was used for identifying mutations in 284 genes, including 118 human genes, three human microRNAs, and 163 human orthologues of mouse deafness genes. The Agilent SureSelect Target Enrichment system was used for a final capture design targeting 4,475 exons of 1.86Mb. The multiplexed libraries, representing 96 Israeli Jewish and Palestinian Arab patients, were analyzed with paired-end sequencing at a read length of 2x101 bp, using the Illumina HiSeq 2000. Coordination with homozygosity mapping in consanguineous families optimized bioinformatics analysis.

**Results**

This method resulted in doubling the number of deafness genes in the Middle East population. Protein structure predictions were made to provide insight into deafness mechanisms. Mutations and the genes identified will be presented.

**Conclusion**

The discovery of novel genes will have implications not only for the Middle East, but worldwide, as many mutations first found in this region have turned out to be present in other populations. This strategy allows for improved diagnostics, in an economically and temporally feasible manner, and establishing etiologically-based genetic counseling and hearing loss management. Further gene discovery will allow for a better understanding of the mechanisms of deafness, facilitating therapeutic development.

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**Wbp2-deficient mice show a progressive pattern of high-frequency hearing loss**

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**Background**

The WBP2 gene encodes the WW domain-binding protein 2. WBP2 is phosphorylated before translocating into the nucleus, where it specifically binds to the estrogen receptor α, working as a transcriptional coactivator (Dhananjayan et al., 2006). Wbp2 knockout mice (Wbp2−/−) were screened for hearing impairment as part of the Wellcome Trust Sanger Institute Mouse Genetics Programme.

**Methods**

ABRs were recorded from ketamine / xylazine anaesthetised mice, aged 4, 14, 28 and 44 weeks. Immunohistochemistry on paraffin sections from five days and four weeks old mice was performed to study Wbp2 expression in the inner ear. Middle ear dissection, inner ear clearing and scanning electron microscopy were performed on four and thirty weeks old mice to characterize the gross structure of the inner ear. For the innervation study, we are using confocal imaging of the sensory epithelium in Wbp2KO/KO and littermate controls at four weeks, using antibodies to CtBP2 to label pre-synaptic ribbons, to GluR2 and GluR3 AMPA subunits to label post-synaptic densities and to Neurofilament to label unmyelinated nerve fibers in the sensory epithelium.

**Results**

Auditory Brainstem Response measurements in Wbp2-deficient mice showed high-frequency hearing impairment as early as 4 weeks old at frequencies from 24 kHz and above. This hearing loss was progressive, with these thresholds becoming higher by 14 weeks and hearing impairment becoming increasingly pronounced at lower frequencies (down to 12kHz) from 28 to 44 weeks. Immunohistochemistry detected Wbp2 expression in inner and outer hair cells of the organ of Corti and in cells of the stria vascularis, spiral ligament and spiral ganglion in control mice. No anomalies were found in the gross structure of the inner ear and the middle ear of Wbp2KO/KO compared to controls, and Scanning Electron Microscopy showed no sign of hair cell degeneration in the organ of Corti.

We are investigating whether the Wbp2KO/KO high frequency loss is due to a problem in the innervation to the basal turn of the cochlea that normally responds best to high frequencies. Preliminary data show swelling of afferent nerve terminals contacting the IHCs as well as an abnormal organisation of ribbon synapses at the inner hair cell membrane in Wbp2KO/KO, and this phenotype is more pronounced in the basal region of the cochlea.

**Conclusion**

These experiments will help us to understand the role of Wbp2 in ear function and to gain more insights on progressive age-related hearing loss.

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**Mislocalization of POU3F4 in the Nucleus due to Novel Mutations in Humans and Mice**

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**Background**

POU3F4 is a POU domain transcription factor that is required for hearing. In the ear, POU3F4 is essential for the mesenchymal remodeling of the bony labyrinth, and is the causative gene for DFNX2 human deafness. Ear abnormalities underlie this form of deafness and include hypoplasia of cochlear regions, reduced coiling of the cochlea, and stapes malformations. A male child of Israeli Jewish Ashkenazi descent presented with congenital profound bilateral hearing loss with a Mondini malformation, one of the hallmarks of X-linked POU3F4 deafness. An ENU screen led to the recovery of a mutant mouse line, schwindel (sdl), exhibiting deafness and circling. Both were further studied to identify the causative mutation responsible for their hearing loss.

**Methods**

The patient's DNA was included in a targeted capture and exome sequencing study. Exons of 284 human genes were captured and sequenced using a 101-bp paired-end recipe to sequence 48 captured library/human DNA samples multiplexed on one lane on an Illumina HiSeq 2000 Analyzer. SNP, indel, and CNV calls were made for all samples. The founder mouse sdl was generated in a large-scale ENU mutagenesis program. After defining linkage to chromosome X using microsatellite markers, Sanger sequencing was used to sequence the candidate Pou3f4 gene. Inner ears were
analyzed by paint-fill, histology, scanning electron microscopy, and immunohistochemistry. COS-7 cells were transfected with the mouse mutation.

Results
Targeted capture and massively parallel sequencing identified a nonsense mutation in the human POU3F4 gene in the child. Sanger sequencing confirmed the presence of c.C235T, predicted to lead to a stop codon, p.Q79X. Analysis of the mouse colony mating data provided evidence that the sdl phenotype is linked to the X chromosome, confirmed in a genome scan. Sequencing of the Pou3f4 gene revealed c.T900A, predicted to cause a stop codon, p.C300X. The subcellular localization of the mouse mutation is damaged and the protein does not appear to be in the nucleus, as seen in transfected cells and Pou3f4\textsuperscript{sdl/sdl} ears.

Conclusion
We report novel POU3F4 mutations in human and mouse. Massively parallel sequencing is shown to be a valid approach for identifying a mutation in a single affected child, with no family history. This approach will continue to speed up the discovery of genes and mutations involved in inner ear pathologies.

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Gene Discovery Approaches to Characterize Molecular Mechanisms Underlying the Brn4/Pou3f4 Mutant Phenotype

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Background
Targeted mutagenesis of the mouse Brn4/Pou3f4 gene generates an animal model for X linked deafness, DFNX2, that accurately recapitulates much of the phenotype observed in DFNX2 patients, including enlarged internal auditory meatus, hypoplastic cochlea and auditory nerve anomalies (Phippard et al., J. Neurosci. 19:5980, 1999; Coate et al., Neuron, 73:49, 2012). Further characterization of the mouse mutant demonstrates that much of the Brn4 phenotype results from dysmorphogenesis during inner ear development.

Methods
To assess the molecular changes that give rise to the mutant phenotype, we have undertaken expression profiling using microarray technology.

Results
We and our collaborators have characterized the molecular mechanisms underlying the enlarged internal auditory meatus (Ahn et al., ARO, 2009) and the auditory nerve anomalies (Coate et al., 2012). Additional gene discovery approaches are directed towards uncovering the molecular mechanisms regulating other aspects of inner ear development that are disrupted in the Brn4 mutants. We will present our analyses of genes whose expression is altered in the mutants, as well as a more in depth analysis of the changes in the TFGß signaling pathway that affect temporal bone development.

Conclusion
These gene discovery approaches should provide additional insights into the underlying molecular mechanisms that direct the development of the inner ear.

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EVALUATION OF THE PATHOGENICITY OF GJB3 AND GJB6 VARIANTS ASSOCIATED WITH NONSYNDROMIC HEARING LOSS

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Background
A number of genes responsible for hearing loss (HL) are related to ion recycling and homeostasis in the inner ear. Connexins (Cx26, Cx31 and Cx30, respectively encoded by the GJB2, GJB3 and GJB6 genes) are core components of gap junctions (GJs) in the inner ear. GJs are intercellular communication channels and an important factor that are
associated with HL. To date, a molecular genetics study of GJB3 and GJB6 as a causative gene for hearing loss has not been performed in Korea. This study was therefore performed to elucidate the genetic characteristics of Korean patients with nonsyndromic sensorineural hearing loss and to determine the pathological mechanism of HL by checking for intercellular communication function of Cx30 and Cx31 variants.

Methods
The GJB3 and GJB6 coding regions were amplified by polymerase chain reaction. An ABI 3130XL DNA sequencer was used to assay the products. GJ biochemical coupling was measured using a single cell dye transfer assay. One of the two adjoining cells forming a GJ was performed patch-clamp using a micro electrode filled with a 1% Lucifer Yellow solution or with 0.15 mM Propidium Iodide. GJ ionic coupling was measured using a Ca2+ transfer assay. Transfected cells grown on cover glasses were loaded with the calcium indicator dye, fura-2 acetoxymethyl ester, and pluronic F-127. Mechanical stimulation was performed by lightly touching one cell membrane, and calcium image was analyzed using Axon Imaging Workbench software.

Results
Sequencing analysis of the GJB3 and GJB6 genes in our population revealed a total of nine variants, including four novel variants in the two genes. Three of the novel variants (Cx31-p.V27M, Cx31-p.V43M and Cx30-p.I248V) and two previously reported variants (Cx31-p.V84I and Cx30-p.A40V) were selected for functional studies using pathogenicity prediction program and assessing for whether the mutation was located in a conserved region of the protein. The results of biochemical and ionic coupling tests showed that both the Cx31-p.V27M and Cx31-p.V84I variants did not function normally when each was expressed as a heterozygote with the wild-type Cx31.

Conclusion
This study demonstrated that two variants of Cx31 were the pathogenic mutations with dominant negative effect. This information will be valuable in understanding the pathogenic role of GJB3 and GJB6 mutations associated with HL.
Gap junction mediated miRNA intercellular communication
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Background
Gap junction is an intercellular channel and possesses a relatively large pore size (1.0-1.5 nm) allowing passage of ions, cell signaling molecules, and small molecules up to 1.5 kDa. miRNAs are small in size (<1.0 nm in diameter and 3.5-8 nm in length) and are able to survive and function for several hours and even days. In this study, we investigated whether miRNA can pass through gap junction to function.

Methods
Connexin-defined cell lines, fluorescent-activated cell sorting (FACS), quantitative real-time PCR (qRT-PCR), and scrape-loading were used to measure miRNA intercellular transfer.

Results
We found that miRNA had transfer between cells in the connexin-defined cell lines but had no intercellular transfer in the connexin-null HeLa cell line. This intercellular transfer of miRNA was eliminated by gap junction channel blockers. Connexin mutations also disrupted such miRNA intercellular transfer. We also used scrape-loading of fluorescence-tagged miRNA to directly show miRNA intercellular passage through gap junctions. miRNA had good intercellular diffusion in the connexin cell lines but had no diffusion in connexin-null cell line and Cx26R75W mutant cell line. The diffusion was also blocked by gap junction channel blockers. The functional assay by miRNA reporter further shows that the transferred miRNA in neighboring cells is functional.

Conclusion
miRNA can pass through gap junctions directly. These studies also indicate that gap junction channels not only play an important role in metabolic communications but also in genetic communications between cells.

A Japanese Family with Autosomal Dominant Auditory Neuropathy Spectrum Disorder
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Background
Auditory neuropathy spectrum disorder (ANSD) is a disorder where outer hair cell function shows normal as revealed by otoacoustic emissions (OAEs) and/or electrocochleography, but auditory brainstem responses (ABRs) is absent or severely impaired. Although the etiology of ANSD is extremely variable, approximately 40% of cases have a genetic background. Non-syndromic ANSD most commonly occurs as a sporadic or recessive pattern. In the present study, we report the clinical features of a Japanese family with autosomal dominant ANSD and the result of next-generation sequencing for the proband.

Methods
A three-generation Japanese family with autosomal dominant sensorineural hearing loss (SNHL) was studied. In 4 of the 10 family members, audiometric examinations including pure-tone audiometry, distortion product OAEs (DPOAEs), and ABRs were performed. For the measurement of ABRs, alternatively polarity clicks at the intensity of 90 and 105 dB nHL
were presented as the acoustic stimuli. Whole exome sequencing was performed on the proband using the Agilent SureSelectXT Human All Exon V4+UTRs+LincRNA and HiSeq 1000 system (Illmina).

Results
The proband was a 29-year-old female who had suffered from gradually progressive, bilateral hearing loss (HL) since the age of 20. Pure-tone audiogram showed bilateral moderate SNHL with normal DPOAEs and absent ABRs. Speech discrimination scores were 90% in the right ear and 70% in the left ear. The proband’s sister (25-year-old) and mother (48-year-old) became aware of HL at the ages of 20 and 32, respectively. Audiograms showed bilateral moderate SNHL in the sister but bilateral profound SNHL in the mother. The sister showed normal DPOAEs but the mother had decreased DPOAEs. Both the two members showed absent ABRs. The proband’s grandmother (71-year-old) had been aware of slight HL during the last few years. She had bilateral mild SNHL, decreased DPOAE, but normal ABRs, suggesting presbycusis. The proband’s grandfather who died at the age of 62 had bilateral profound HL by the age of 60. All the affected members had no symptoms other than HL. The result of next-generation sequencing for the proband showed no variants in OTOF, PJVK and DIAPH3 genes.

Conclusion
A Japanese family with late onset, gradually progressive, autosomal dominant ANSD was shown. Further genetic studies for the other family members are needed to know the etiology of the ANSD.

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Environmental and lifestyle factors involved in normal hearing function and age-related hearing loss
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Background
Until now, little is known about environmental/lifestyle factors underlying normal hearing function and age-related hearing loss (ARHL). To reach this goal we decided to run an epidemiological study on hearing quantitative and qualitative traits (analysing the low, medium and high Pure Tone Average-PTA) on subjects coming from several isolated populations from Europe, Caucasus and Central Asia.

Methods
4401 subjects coming from Europe (Italy, Croatia and Ukraine), Caucasus (Armenia, Azerbaijan, Georgia) and Central Asia (Uzbekistan, Kazakhstan, Tajikistan and Kirghizstan) and ranging from 4 to 95 years were included in the analysis. Different thresholds (0.25, 0.5, 1, 2, 4, 8 kHz) as well as PTAs have been considered in the study. A medical examination has been performed in order to exclude pathological conditions related to the hearing impairment. A series of lifestyle/environmental factors have been tested including habits (smoking, drinking, etc.), diet (several foods and beverages intakes), level of education, etc. A model based on linear/logistic regression was used to fit PTAs to all the covariates (smoking, chocolate, coffee, tea, wine, beer, dairy products, hard-liquor) including sex and age.

Results
As regards hearing function, our results show, for the first time, that among eight analyzed variables only coffee consumption (but not the coffee intake) was strongly associated with better normal hearing function in particular at low and high frequencies in four out of ten countries investigated (p=0.038 in Sardinia, 0.006 in Southern Italy, 0.017 in Azerbaijan, 0.016 in Tajikistan) as well as PTAs have been considered in the study. A medical examination has been performed in order to exclude pathological conditions related to the hearing impairment. A series of lifestyle/environmental factors have been tested including habits (smoking, drinking, etc.), diet (several foods and beverages intakes), level of education, etc. A model based on linear/logistic regression was used to fit PTAs to all the covariates (smoking, chocolate, coffee, tea, wine, beer, dairy products, hard-liquor) including sex and age.

Conclusion
As regards ARHL phenotype, we found an association in two Italian isolated populations (p=0.02 for Sardinia, and p=0.05 for Southern Italy), in which a lower level of education is associated to an increased susceptibility to ARHL (Figure 2).

These data demonstrate, for the first time, a relationship between coffee consumption and a better hearing function, and confirm the relevance of a higher education in protecting against ARHL. The understanding of lifestyle/environmental risk factors will help us in developing new preventive strategies for ARHL.
Linkage studies and Whole Exome Sequencing analysis aimed at the identification of new deafness genes

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Background
Nonsyndromic Hereditary Hearing loss is a common disorder accounting for at least 60% of prelingual deafness. Despite GJB2, GJB6 and A1555G mitochondrial mutations play a major role worldwide still there is a need to search for new causative mutations/gene underlying the disease in several populations such as Italy as well as countries from the Gulf region

Methods
A combined strategy has been developed based on linkage analysis and whole exome sequencing. In particular, high density SNPs arrays (OmniExpress 700K from Illumina, USA) have been utilized to get linkage data (even the so called “sniff” considering that families in some cases are not large enough to lead to the identification of only one locus). By this approach it is possible to define a given number of candidate loci for each family to be applied in the exome sequencing filtering phase. As regards exome sequencing they have been carried both on SOLID4 or Illumina platforms after the enrichment step carried out using Agilent technology. After library construction and sample running, single nucleotide variants (SNVs) were called by Samtools V0.1.8 and filtered comparing with dbSNP database v132 and our in-house database. Selected variants were then confirmed by direct Sanger sequencing and a complete functional prediction analysis was done using a series of tools such as PolyPhen-2, MutationTester, SIFT, Consurf, and Condel

Results
Six Italian families (showing a dominant pattern of inheritance) and 5 Qatari ones (showing a recessive pattern of inheritance), all negatives for the presence of mutations in the most common HHL genes (i.e. GJB2, GJB6 and A1555G mitochondrial mutation) have been selected. In the Italian families our approach led to the identification of 5 new loci/gens for which Sanger sequencing confirmation is now in progress. Moreover, 4 new loci and one new possible HHL gene have been identified in Qatari families. In particular a causative mutation (c.7873 t>g leading to p.*2625Gluext*11) in BDP1 gene has been found in one family. The mutation disrupts the termination codon of the transcript resulting in an elongation of 11 residues of the BDP1 protein. Immunohistochemistry analysis carried out in the mouse inner ear showed Bdp1 expression in the mouse cochlea

Conclusion
These findings definitely increase our knowledge of new HHL genes, further confirming the importance of the combination of both strategies for disease gene identification and may suggest new targets for hearing impairment treatment and prevention
Auditory Rehabilitation and Genetic Analysis in Patients with BO/BOR Syndrome Showing Mixed Hearing Loss

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Background
BO/BOR syndrome is one of the most common forms of autosomal dominant syndromic hearing loss. The EYA1 gene has been identified as the causative gene of BO/BOR syndrome, and association with mutations in SIX1 and SIX5 genes has also been reported. In this study, we have focused on the auditory manifestations and rehabilitation in patients with BO/BOR syndrome, and also performed genetic analysis for EYA1, SIX1, and SIX5 genes.

Methods
Seven families (10 patients) including one multiplex family showing hearing loss and one or more of the typical features of BOR syndrome were included. Pure tone audiometry was serially performed, and the type, onset, progressiveness of hearing loss were evaluated in each patient. The history and outcome of middle ear surgeries, hearing aids, and cochlear implantation were analyzed. The temporal bone CT and temporal MRI were analyzed. For genetic analysis, direct sequencing and deletion analysis of the EYA1 gene were performed. If mutation was not identified in the EYA1 gene, mutations in the SIX1 and SIX5 genes were subsequently analyzed. For splice site mutations, exon trapping analysis was performed.

Results
All patients presented with mixed type of hearing loss ranging from moderate to profound degree. The onset was mostly perilingual or postlingual, and five of 10 patients presented variable degrees of progressive hearing loss. All patients exhibited enlarged vestibular aqueduct of tubular type, cochlear hypoplasia, facial nerve anomaly, and ossicular abnormalities on temporal bone CT. Five of 10 patients underwent middle ear surgeries after which none of the patients experienced significant hearing gain. All of the patients used hearing aids. Cochlear implantation was performed in two patients with severe to profound hearing loss, which improved their hearing and language ability significantly. Five of 7 families presented with mutations in the EYA1 gene including 3 splice-site and 1 missense mutations as well as a whole gene deletion found in one patient. No significant genotype-phenotype correlation was demonstrated concerning auditory manifestations. No mutations in the SIX1 or SIX5 gene were found in patients without EYA1 gene mutation.

Conclusion
Four novel mutations of the EYA1 gene and one whole gene deletion were identified in patients with BO/BOR syndrome showing mixed hearing loss and inner ear anomalies. Various auditory manifestations were demonstrated but no genotype-phenotype correlation could be found. Middle ear surgeries failed to improve hearing in these patients, while cochlear implantation provided successful auditory outcome in selected patients.

Functional Studies of the Mouse Deafness Gene Grxcr2

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Background
Mice carrying a targeted mutation of Grxcr2 exhibit severe hearing loss that is associated with developmental defects in stereocilia bundle orientation and organization, similar to those previously observed in mouse models of Usher Syndrome. The localization of GRXCR2 protein within stereocilia suggests that it plays an intrinsic role in influencing normal orientation, organization and cohesion of the bundles. Based on sequence analysis suggesting a putative chaperone activity of GRXCR2, together with the similarity in bundle defects observed in mouse Usher and Grxcr2 mutants, we hypothesize that GRXCR2 normally acts to assist in appropriate folding and/or complex formation of Usher proteins and thereby regulates their localization and function.

Methods
Predicted protein structure of GRXCR2 was evaluated using the Phyre2 (Nature Protocols 4:363-371, 2009) and I-TASSER (Nature Protocols 5:725-738, 2010) platforms. Localization of Usher proteins was determined by immunocytochemistry of fixed cochlear tissue derived from Grxcr2 mutants and controls using specific antibodies.

Results
We identified significant similarity in the predicted structure of the Cys-rich, C-terminal region of GRXCR2 and the known secondary structures of several dnaJ-related proteins, including E. coli dnaJ/Hsp40 and human DNAJ3. This family of

In early postnatal Grxcr2 mutants, we observed a broad stereocilia distribution of the Usher protein harmonin rather than the precise tip localization found in bundles of normal control mice. This is unlikely to be a general effect on stereocilia proteins, as in previous studies we have demonstrated intact transduction currents in OHC bundles in early postnatal Grxcr2 mutants, indicating that assembly of the apparatus necessary for mechanotransduction such as tip links initiates normally.

Conclusion
The similarity of the C-terminal domain of GRXCR2 with Cys-rich domains present in dnaJ-related co-chaperone proteins suggests a role in regulating protein folding and/or complex formation in stereocilia. Consistent with this model, we have observed mislocalization of harmonin early in bundle development in Grxcr2 mutant mice. We are currently testing for direct interactions of GRXCR2 with harmonin and the localization of other Usher proteins in mutant stereocilia.

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Intra-familial Locus Heterogeneity Confounds Deafness Gene Identification

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Background
Pedigree-based linkage analysis and homozygosity mapping have been used to genetically map the chromosomal locations of human hereditary recessive deafness loci in consanguineous pedigrees and community isolates. Recessive nonsyndromic deafness is both highly genetically heterogeneous and not rare. Although most, if not all, DFNB loci have been identified in pedigrees where deafness segregates as a monogenic trait, the population incidence and extreme heterogeneity suggest that some pedigrees may be segregating mutant alleles of more than one deafness gene. Pedigrees that fail to yield statistically significant evidence of genetic linkage may do so because two or more non-allelic pathogenic mutations are segregating within the same pedigree.

Methods
We reanalyzed consanguineous pedigrees from Pakistan in which recessive nonsyndromic deafness was segregating in multiple sibships. Each of these pedigrees failed to show evidence of linkage to a single locus in genome-wide screens with 388 STR markers. To address the possibility of locus heterogeneity, we Sanger-sequenced several known deafness genes in at least one affected individual from each sibship. Also, in a large North American kindred with multiple sibships segregating deafness, we used Sanger-sequencing, genotyping by restriction enzyme assay, or array CGH to identify multiple non-allelic mutations.

Results
Intra-familial locus heterogeneity was documented in six deafness pedigrees. In five of the pedigrees, two non-allelic pathogenic mutations are segregating. All affected siblings in each sibship are homozygous by virtue of identity by descent for one of these two mutations; however, closely related sibships in each pedigree are segregating pathogenic alleles of different genes. In each pedigree, homozygous pathogenic alleles of GJB2 are responsible for deafness of some individuals while other deaf individuals in the same pedigree are homozygous for pathogenic alleles of SLC26A4, HGF, or MYO7A. In the North American pedigree, two recessive DFN1 mutations, two recessive mutations of SLC26A4, and a novel deletion of POU3F4 segregate in this community isolate.

Conclusion
Intra-familial locus heterogeneity in hearing loss is uncommon or perhaps largely unpublished, but should no longer be surprising, because congenital hearing loss is not rare, a significant proportion is genetic, and many genes cause monogenic hearing loss. Unrecognized locus heterogeneity can confound deafness gene identification by linkage analysis or homozygosity mapping when all affected and unaffected individuals in different sibships are included in the analysis. As whole-exome sequencing becomes established in clinical practice, it is important to consider the potential for genetic heterogeneity in extended families.
Foxo3 Protects Hearing Through Synapse Maintenance
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Background
Foxo3 is a member of the Foxo winged-helix transcription factor family. Foxo3 mediates oxidative stress responses, apoptosis, and cell cycle withdrawal in many systems. Foxo3 can be activated by excitotoxicity or mechanical stress. Human Foxo3 alleles are associated with extreme longevity and with female infertility. We have discovered that Foxo3 is expressed in cochlear cells, and we hypothesize that it functions to preserve hearing. Previously, we showed that loss of Foxo3 causes high-frequency hearing dysfunction with age, and here we have characterized the mechanism of this hearing loss.

Methods
All experiments were conducted on Foxo3 wild-type and knock-out littermates, derived from FVB/n, at two and four months of age. Immunohistochemistry was used to analyze Foxo3 expression in the sensory region of the ear and to detect hair cells and neurons for subsequent quantification. Auditory function was assessed with Auditory Brainstem Responses (ABR) and Distortion Product Otoacoustic Emissions (DPOAE). Ultrastructural analysis was performed on the 32-kHz cochlear turn isolated from both wild-type and Foxo3-KO animals at four months of age.

Results
Foxo3-KO mice develop hearing loss at 24 and 32 kHz by four months of age, compared to their wild-type counterparts. Outer hair cell function, assessed by DPOAE amplitudes at the same frequencies, appears unaffected. Foxo3-KOs also have similar numbers of hair cells compared to wild-type littermates. Moreover, spiral ganglion neurons are still present in the Foxo3-KO in the 32-kHz cochlear turn. We find, however, that synapses between spiral ganglion neurons and inner hair cells are dysmorphic, with increased gaps, reduced post-synaptic densities and neurite beading.

Conclusion
We conclude that Foxo3 preserves hearing with age by maintaining synapse integrity. We speculate that the synapse disruption exhibited in the four month-old Foxo3-KO mouse is due to an intrinsic sensitivity to excitotoxicity. Future research will address the mechanism of Foxo3’s role in noise damage to the cochlea.

Mapping of an otitis media gene within an indigenous population
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Background
Otitis media (OM) affects a large proportion of the global population, causing significant mortality and morbidity, and is a major cause of hearing loss across all age groups. It incurs a huge cost to health systems from repeated consults and antibiotic prescription. Despite vaccination regimens and specific treatment guidelines for different forms of OM, due to changes in pathogen patterns and increased antibiotic resistance, OM remains an important public health problem. Chronic OM or recurrent/persistent OM/middle ear effusion may require surgical intervention and can result in permanent hearing impairment which influences a person’s communicative skills, cognitive abilities and psychosocial well-being. Strong evidence for a genetic basis of OM susceptibility exists, but only a few genes have been associated with OM in humans, and the causal variants are yet to be identified.

Methods
A single large pedigree from an indigenous, intermarried population with a high prevalence of OM has been ascertained. Out of 49 pedigree members who provided DNA, 42 DNA samples were successfully genotyped using the exome chip. Minimal pedigree errors were detected using PedCheck for Mendelian inconsistencies and MERLIN for possible genotyping errors. Two-point and multipoint linkage analyses were performed using Superlink, and haplotypes were reconstructed using Allegro software.

Results
For the initial linkage analyses, we used 7,601 common variants from the exome chip which were informative for the pedigree. Based on an autosomal dominant model of inheritance with reduced penetrance (70%), the maximum two-point LOD score for the entire pedigree was 1.54 at chromosome 5p, and 1.43 at the 10q region. When affecteds-only analysis was performed at equal allele frequencies, the maximum LOD scores slightly increased to 1.58 at 5p and 1.56 at 10q. For
multipoint analysis, rare variants within the 5p and 10q regions that are informative for the pedigree were included. Using affecteds-only, the maximum multipoint LOD score at 5p is lower at 2.77, while at the 10q region the maximum multipoint LOD score is statistically significant at 3.56. Linkage heterogeneity within the pedigree was detected.

**Conclusion**

DNA samples from affected individuals have been submitted for exome sequencing, which will lead us to identification of genetic variants that cause OM susceptibility. The identification of genetic variants that confer susceptibility to OM has the potential to increase our understanding of pathophysiology, which can then lead to improvements in preventive and therapeutic strategies for OM.

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**Cell Type-Specific Studies of the Mouse Inner Ear – Challenges and Solutions**

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**Background**

The auditory and vestibular sensory organs consist of complex epithelia that are composed of multiple cell types. Fluorescent Activated Cell Sorting (FACS) has become an increasingly popular tool to study the development of the mouse inner ear, isolate specific cell populations for subsequent applications, or study the downstream effects of cell type-specific mutations. In this presentation we describe OTO-SORT, a protocol that can be used to isolate the sensory epithelium, neurons, vascular endothelium and mesenchyme from the auditory and vestibular systems, as well as combinations of OTO-SORT with transgenic mice expressing cell type-specific fluorescent proteins. We address the technical aspects of the process in a stepwise fashion, starting from the cell dissociation to challenges and solutions for downstream applications.

**Methods**

Tissue is dissected from mice up to postnatal day 5, incubated in thermolysin and then dissociated in Accutase enzyme cell detachment medium, followed by a mechanical disruption using a 23G blunt ended needle connected to a 1ml syringe. The reaction is stopped by adding an equal volume of complete media. Cells are passed through a 40-µm cell strainer and washed in FACS buffer. The dissociated cells are then stained with CD326-APC, CD49f-alexa488 and CD34-PE prior to sorting.

**Results**

The auditory and vestibular epithelia can be dissociated with minimal induction of cell death. As little as one ear can be used for isolation of hair cells, for example, with a yield that can reach up to 1000 cells per auditory or vestibular epithelium. Tissue can be dissected and successfully sorted up to 2 days following dissection. Cell type-specific gene expression analysis can identify transcription factors and miRNAs that determine cell fate.

**Conclusion**

FACS can be applied to study inner ear development in embryonic and early postnatal stages even when starting from single epithelia. Using a combination of antibodies for cell surface markers and mice that express fluorescent molecules in a cell type-specific pattern can enhance the ability to sort unique cell populations. Generating antibodies for the extracellular domain of cell type-specific transmembrane proteins will simplify FACS of the inner ear and obviate the need to use mice expressing cell type-specific markers.
Molecular Network Analysis of Hearing Loss and Hypogonadism

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Background
Hypogonadism and hearing loss are conditions that have significant clinical burdens. They represent a diverse range of clinical presentations and pathological processes. While they are both functionally distinct entities, the gonadal axis and auditory pathway have similar embryological origins. Syndromic forms of hypogonadism with hearing loss are well documented. However, identification of specific genes involved in both non-syndromic hearing loss and hypogonadism have yet to be elucidated. The aim of this study is to identify key molecular regulators of hypogonadism and hearing loss through a comprehensive bioinformatic analysis of genes common to both diseases.

Methods
A search of the Online Mendelian Inheritance in Man (OMIM) database was performed to identify genes associated with hypogonadism. All entries including the term hearing loss were compiled into a dataset. Two network analyses were performed using Ingenuity Pathway Analysis software (IPA; Ingenuity Systems, Redwood City, California) – one focused on direct connections among genes, and another, inclusive of direct and indirect connection. The most interconnected, i.e. nodal molecules, of each networks were identified. Bioinformatic results were validated by immunohistochemistry applied to cross-sections from human cochleae and gonadal tissue (male and female).

Results
The OMIM search identified a total of 59 genes associated with both hypogonadism and hearing loss. The IPA analyses yielded five highly statistically significant networks. The transcription factor hepatocyte nuclear factor-4α (HNF4α) emerged as a key nodal molecule in both networks. This bioinformatic result was validated by identifying, via immunohistochemistry, specific expression of HNF4α in the human reproductive tract and inner ear.

Conclusion
This study identifies key nodal molecules that are likely to play significant roles in both hypogonadism and hearing loss. We highlight HNF4α; as a potential novel regulator of both the gonadal axis and the auditory system. The precise role of HNF4α; in these tissues requires further study.

Acoustic Trauma Changes the Expression of Serotonergic Receptors in the Inferior Colliculus

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Background
Multiple studies have demonstrated that an important neuromodulatory network in the inferior colliculus (IC) is the serotonergic system, which modulates neural activity in response to acoustic stimuli. In this work, we studied the effects of acoustic trauma on the expression of serotonergic and GABAergic receptor genes in order to determine how acoustic insult to the inferior colliculus impacts the neuromodulatory network.

Methods
Individual male mice (CBA/J strain) at 11-15 weeks of age were used for these experiments. The auditory brainstem response (ABR) was measured at four frequencies (8, 12, 16, and 20 kHz) to establish hearing thresholds. Sham surgical controls were then left under anesthesia for three hours while trauma individuals were anesthetized and exposed to a 10 kHz pure tone at 116 dB for three hours. After a one month recovery period, the auditory brainstem response was measured again to determine hearing loss and the individual was sacrificed. The IC was then dissected and processed for real-time quantitative PCR (RT-qPCR). The expression of the following genes was measured relative to β-actin: 5Htr1B, 5Htr1A, 5Htr2A, 5Htr3A, and GABArA.

Results
ABR measures indicated that the trauma paradigm used caused severe hearing loss at frequencies above the trauma stimulus in individuals after a month of recovery time (threshold shifts of 40dB or greater at 12, 16 and 20 kHz). Significant changes between sham surgical controls and traumatized individuals were found in the expression of two genes following acoustic trauma. As other researchers have reported, expression of the GABArA gene was downregulated 60% on average. In contrast, the 5Htr1B gene was upregulated 3-6 fold in traumatized individuals. This results in a mean 4.6-fold
relative shift of GABA to 5Htr1B between control and trauma individuals. The other genes measured were not significantly different between sham controls and trauma.

**Conclusion**

Since the 5-HT1B receptor may act in the IC by decreasing GABA release, our gene expression results suggest that increased 5-Ht1B expression contributes to a general downregulation of inhibitory mechanisms following acoustic trauma.

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### Conditional expression of microbial rhodopsins in mouse cochlear hair cells and supporting cells

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Microbial rhodopsins are light-gated ion channels that can be expressed in excitable mammalian cells to modify their activity. ChR2(H134R) is a channel rhodopsin variant that produces reversible membrane depolarization due to cation influx in presence of blue light and can activate neurons. Archaerhodopsin-3, an outward proton pump, and Halorhodopsin, an inward chloride pump, under yellow light produce membrane hyperpolarization and neuronal silencing. Recently, the Allen Institute for Brain Sciences has created floxed mice of these three variants. Cre-mediated deletion of a stop cassette in these mice produces expression in specific neuronal populations and successful modulation of their activity. We have crossed these floxed rhodopsin mice with CreER mouse lines that are specifically expressed in different cell types of the postnatal cochlea. PrestinCreERT2 and Fgfr3iCreER drive specific expression of rhodopsins in outer hair cells and supporting cells, respectively, when induced with tamoxifen (3mg/40g bodyweight) once a day at postnatal day (P) 6 and P7. Auditory brainstem responses measured in these mice show that hearing is largely not affected by microbial rhodopsin expression in cochlear cells. The models driven by PrestinCreERT2 can be used to alter (silence or augment) electromotility of isolated outer hair cells with light, overriding the membrane potential changes that trigger this response. In vivo, light stimulation may be used to modulate outer hair cell-dependent amplification at specific locations along the cochlea to determine the place of generation of the peak of sound-driven travelling waves. Supported by: Wellcome Trust Fellowship (MML), NIH, ALSAC and Hartwell Foundation.

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### In vivo visualization of endolymphatic hydrops using optical coherence tomography (OCT)

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**Background**

Endolymphatic hydrops has been believed as the most important feature for the pathogenesis of Meniere’s disease. However, its dynamic pathology is not well understood due to the lack of in vivo non-invasive testing tools for the inside morphology of inner ears in animals and humans.

Optical Coherence Tomography (OCT) is a relatively new imaging modality that utilizes near infra-red light to obtain cross sectional images of opaque biological tissues. OCT has already been used as a diagnostic imaging tool in ophthalmology and coronary vessels.

In this study, we demonstrate that OCT is capable to visualize inner ear structure of endolymphatic hydrops model mice in vivo, and discuss the possibility of OCT as an in vivo diagnostic tool for inner ear disorders.

**Methods**

Eleven week-old mice with targeted disruption of *Slc26a4* (formerly known as Pendrin or Pds) and their littermates were used as experimental animals, because homozygotes for that gene are known to develop extensive dilatation of the scala media (Everett, 2001; Wangemann, 2009). To test the hearing function, auditory brainstem responses (ABRs) were recorded at 8, 16, 32 kHz. On the following day, each animal was anesthetized and the left bulla was removed to expose bony contour of the left cochlea. The animal was placed to obtain the optical sections that include the apex and the round window niche, then a series of cross sectional images was obtained by the OCT (OCS-1300SS, Thorlabs Inc., NJ). Then, animals were intracardially perfused with PBS followed by 4% paraformaldehyde (PFA), temporal bones were excised, inner ears were re-fixed by cochlear perfusion, and postfixed for more than 4 h. Hematoxylin and eosion staining of 10 µm-thick frozen sections were used for morphological evaluations.

**Results**

Homo mice showed profound hearing impairment, and hetero and wild type (wt) mice were normal in hearing. OCT images of hetero and wt mice cochlea revealed the Reissner’s membrane at normal position. While, OCT images of homo
mice revealed limited number of cochlear turns, and the Reissner’s membrane close to the inner wall of the cochlea indicating severe dilatation of the scala media. These findings were consistent with histological study.

Conclusion
OCT is capable to visualize morphological features in the inner ear of live animals. OCT should provide diagnostic information of inner ear diseases in vivo.

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Evaluation of the internal structure of the normal and pathological guinea pig cochleae using optical coherence tomography
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Background
Structural observation of cochlea has primarily been limited to histological methods, which require chemical fixation followed by dissection of the tissues or embedment in mold such as paraffin and sectioning. It is, however, well known that these preparations of tissue samples introduce significant changes in tissue integrity and organization, which thus may induce misinterpretation, limiting the generality and overall value of the results. To innovate the approaches to visualize the internal structure of the cochlea, an emerging noninvasive imaging modality of optical coherence tomography (OCT) has been applied.

Methods
20 Hartley guinea pigs were used and allocated into the following four groups, each consisting of five animals. 1) Normal control group. 2) Endolymphatic hydrops group: ears with the electro-cauterization of the endolymphatic sac and 4-week feeding. 3) Kanamycin sulfate-ethacrynic acid group: administration in combination of kanamycin sulfate and ethacrynic acid and 2-day feeding. 4) Streptomycin sulfate group: ears with perilymphatic perfusion with 20% streptomycin sulfate and 4-month feeding.

To observe the cochleae with OCT, we obtained the both temporal bones immediately following fixation and kept them in 10% formalin solution for 1 week. Subsequently, the specimens underwent decalcification in EDTA for 14 days. Then we obtained images of the cochleae by using Santec OCT system (Santec Co., Aichi, Japan). After obtaining OCT images, the temporal bones were dehydrated in increasing concentrations of alcohol, embedded in paraffin, and cut serially at 6 μm in the plane parallel to the modiolus. The sections were stained with hematoxylin and eosin and observed under a light microscope.

Results
In this OCT study, we visualized the internal structures of the organ of Corti, Reissner’s membrane, and lateral wall in the normal and pathological ear. OCT images of normal and pathological cochleae did not exhibit artifacts observed in HE specimens that were induced during histological preparations. OCT could demonstrate endolymphatic hydrops, strial atrophy, and the damage of the organ of Corti, shown as the distention of the Reissner’s membrane, thinning of the lateral wall, and flattening of the organ of Corti, respectively.

Conclusion
By decalcifying the bony wall of the cochlea, we could clearly and widely visualize the internal structures of the normal and pathological cochleae. These OCT images did not exhibit artifacts observed in HE specimens. These results indicate that observing the decalcified cochlea by using OCT would be of great value of examining the cochlear pathology, prior to or without histological examination.

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The Limited Use of Micro-CT in Cochlear Imaging of Rat
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Background
After the clinical computed tomography (CT) was first developed in 1967 by G. Hounsfield, x-ray CT has been widely used for diagnosis in biomedical fields. X-ray CT is a medical imaging method using tomography created by digital geometry processing to generate a three dimensional image of the inside of an object. The use of x-ray CT has increased dramatically over the last two decades in many countries and has been applied to various organs such as brain, lungs,
pulmonary angiogram, cardiac, abdominal and pelvic, and extremities. The use of x-ray micro CT on the ear has been mainly focused on the outer and middle ears. The objective of this study is to investigate the degree of visualization for the macro-, micro-, and nano-structures of the cochlea with x-ray micro-CT based on a cone-beam geometry for rats.

Methods
Before obtaining the visual images, the hearing thresholds were obtained with auditory brainstem responses (ABR). The entire experiments for visual images with x-ray micro CT were performed at Wonkwang University. X-ray micro-CT based on a cone-beam geometry for a small animal was used for examining cochlear image of rats. The micro-CT was mainly composed of an x-ray tube with tungsten target, a sample holder made from plastic of 2 mm thick, and a digital detector of CMOS (Complementary Metal Oxide Semiconductor).

Results
X-ray micro CT provided a clear depiction of the basal, middle, and apical cochlear turns. X-ray micro CT allows visualizing the macro-structures of the cochlea. However, it did not well depict the micro- and nano-structures of the cochlea.

Conclusion
X-ray micro CT provided a clear depiction of the basal, middle, and apical cochlear turns. X-ray micro CT allows visualizing the macro-structures of the cochlea. However, it did not well depict the micro- and nano-structures of the cochlea. To visualize the micro- and nano-structures of the cochlea, optimal parameter values of x-ray micro CT should be identified. In addition, many factors such as signal-to-noise-ratio, resolution, image averaging, pixel, pixel number, pixel value, exposure time, and binning should be properly considered.
Two photon microscopy of the mouse cochlea in situ for cellular diagnosis
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Background
Sensorineural hearing loss is the most common type of hearing loss worldwide yet the underlying cause is typically unknown because the inner ear cannot be biopsied today without destroying hearing, and intracochlear cells have not been imaged with resolution sufficient to establish diagnosis. Intracochlear imaging has been technologically challenging because of the cochlea’s small size and encasement in bone.

Methods
Six week old CBA/CaJ mice were exposed to an octave band noise of 8-16 kHz at 106 dB SPL for 2h, which produced permanent cellular damage. Unexposed age- and sex-matched mice served as controls. The cochleae were extracted 3 weeks after noise exposure, and imaged in situ, through the intact round window, after intracaradiac perfusion with 4% paraformaldehyde. A separate group of unfixed cochleae were also imaged. Cochlear imaging was performed with two-photon excitation fluorescence (TPEF) microscopy, one-photon excitation fluorescence (OPEF) microscopy, and wide-field transmission microscopy.

Results
We found that the TPEF yielded clear advantages over OPEF and wide-field transmission microscopy. Our experiments revealed fine cellular structure of the hair cells, and allowed morphological distinction between healthy and noise-damaged organs of Corti. Furthermore, cochlear neurons were imaged through the intact bone. The intensity of TPEF signal was brighter in the freshly harvested, unfixed samples than in the fixed samples. The detected TPEF signal coincided with the peak of emission of flavin adenine dinucleotide (FAD), which is a major endogenous fluoropore in the inner ear. Our results, combined with the fact that the sensory epithelium of the inner ear has one of the highest tissue concentrations of FAD known, strongly motivate future development of TPEF-based microendoscopy for cellular diagnosis of sensorineural hearing loss.

Conclusion
Our results demonstrate that the round window provides a useful access to the cochlea through the middle ear, and they motivate future development of new and efficient diagnostic tool based on two-photon micro-endoscopy. The use of TPEF eliminates the need for exogenous labeling and provides the maximum non-invasiveness in the process.
Variations in microanatomy of the human cochlea
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Background
The human cochlea shows considerable interindividual variability in size and morphology. For development of atraumatic CI electrodes details of variability of human anatomy are required with high precision. For this purpose, X-ray microtomography (µCT) was used. This imaging technique allows for non-invasive high resolution visualization with almost histological quality of the cochlear structure. The obtained data were imaged and dimensions of the scala tympani were determined.

Methods
From fresh frozen temporal bones, blocks of approximately 3, 5 x 3, 5 cm were cut out around the cochlea. In order to visualize the soft tissues in the cochlea, the cochlear fluid was gently removed. The bones were scanned using a Skyscan 1173 machine (40-130kV source, 5Mp, 12 bit CCD sensor), resulting in images with pixel size varying from 8-17 μm. Materialise MIMICS software was used to segment out the scala tympani (ST) and Rosenthal’s canal from the obtained µCT images. A 3-D model of the segmented area was generated. To compare multiple cochlear dimensions, the centers of the round window were merged, and the cochlear coordinate system was applied for all cochleae. Cross sectional images were taken perpendicular to the centerline of the ST. The 2-D images were exported to Matlab and analyzed by a custom-made software.

Results
Comparison of different ST showed a large variability in the cross sectional diameter (CSD), vertical layout, and the width and height of the ST. The CSD of the ST reaches a maximum within the first 2 mm and then overall decreases towards the apex. Relative standard deviation of the CSD are between 7.5 and 16.5 %. In general, the mid-scala and modiolar region of the ST contain largest surface area, laterally, the ST surface area diminishes due to the decrease in height. There was significant variability in wrapping and vertical lay-out between the ST: in Z direction, dips were observed regularly between 95 and 130 degrees. A typical incline was observed between 245 and 285 degrees. The Rosenthal’s canal extended up till 460 to 585 degrees.

Conclusion
We segmented the ST and measured the internal dimensions using micro-computed tomography. We observed a large dimensional variability between different ST. These differences could have large implications in the design approach of CI arrays especially towards their capability to preserve residual hearing during insertion of the CI electrode array.
The retardance of the spiral ligament and the basilar membrane decreases from the cochlear base to the apex, compared to the more uniform retardance of other structures.

Conclusion
qPLM has important advantages over immunohistochemistry in evaluating the organization of collagen fibrils and myelinated neuronal processes because qPLM provides detailed qualitative and quantitative information on unstained fiber networks. Our results strongly motivate the future application of polarization-sensitive optical coherence tomography and similar imaging technologies to the human inner ear in vivo.

Intravital Calcium Imaging of Cochlear Hair Cells
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Background
In vivo imaging has drawn much attention in recent years, as it can elucidate the complex biological and pathological events within a living animal. Although histological examination of excised tissues has long served as a fundamental approach for tissue analysis, intravitral confocal microscopy provides instant histopathology at the cellular and subcellular level. Therefore, it is considered ideal for investigating dynamic events involved. In this study, we examine the development and application of intravitral confocal microscopy for calcium imaging of cochlear hair cells upon sound stimulation.

Methods
Adult guinea pig was anesthetized, and apical turn of the cochlea was surgically exposed via ventral approach, leaving tympanic membrane and auditory ossicles intact. A membrane-permeant green-fluorescent calcium indicator, Fluo-4 AM, was applied for 20 minutes, and then removed to fill the middle ear with air. Fluorescent images were obtained before, during, and after sound stimulation. All picture/movie acquisitions were performed using a Nikon A1R confocal laser scanning microscope system attached to an upright ECLIPSE Ni equipped with a x10, 0.3 numerical aperture, 17.3 mm working distance objective. The A1R incorporates both a conventional galvano scanner and a high-speed resonant scanner together. The resonant scanner allows an acquisition speed of 30 frames per second while maintaining high resolution of 512 x 512 scanned points. Fluo4-AM was excited with a 488 nm argon laser, and emission was detected through a 525/50 nm band pass filter.

Results
Inner hair cells, outer hair cells, and supporting cells were all clearly identified, indicating successful loading of Fluo-4 AM dye. Increased fluorescence intensity of the hair cells was observed upon sound stimulation.

Conclusion
We have demonstrated a technique to capture the image of dynamic fluorescent changes in the sound-stimulated cochlear hair cells in living guinea pigs. Although apical opening is necessary for visualization, the preparation well preserves physiological hearing functions and conditions such as sound conduction, innervation, blood supply, and body temperature. Our technique offers opportunities to investigate hair cell transduction of living and hearing animals in both spatial and temporal resolution.

Phosphatidylcholine species localization in the mouse inner ear by mass microscopy
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Background
We recently developed a mass microscope consisting of a mass spectrometry imager with high spatial resolution equipped with an atmospheric pressure matrix-assisted laser desorption/ionization and quadrupole ion trap time-of-flight
analyzer. Phosphatidylincholine (PC), a phospholipid, is a basic structural component of cell membranes. PC species exhibit various binding patterns with fatty acids and were analyzed in the brain, eye, and others; however, the distributions of PC species in the inner ear have not been fully understood. We have previously reported the localization of PC species in the guinea pig cochlea by this microscope. In this study, we extended the use of our mass microscope to the mouse inner ear with special attention to the vestibular organ.

Methods
Three-week-old male DBA2J mouse were anesthetized with pentobarbital and decapitated in accordance with an animal study protocol approved by the Hamamatsu University School of Medicine Animal Care and Use Committee. The temporal bone was isolated and the inner ear was dissected. The inner ear was immediately frozen in liquid-nitrogen-cooled isopentane without fixation. The frozen section was then immersed in a 2% sodium carboxymethylcellulose solution for embedding and stored at -80°C. Before sectioning, the inner ear was held for 20 min at -20°C. Consecutive 15µm sections of the frozen inner ear was prepared using a cryostat. The section was mounted on indium-tin-oxide (ITO)-coated glass slides.

A mass microscope (Shimadzu Corporation, Kyoto, Japan) is a high-resolution IMS instrument with MALDI-QIT-TOF combined with an optical microscope under atmospheric pressure. Matrix was prepared 50 mg/ml 2,5-Dihydroxybenzoic acid (DHB). After the DHB matrix was allowed to dry, the ITO-coated glass slide was placed on the mass microscope target plate. The mass microscope was operated in the range of m/z 400 - 1000 to detect the ion signals of the PC species.

Results
The mouse specimens are smaller than those of the guinea pig, we clearly localized some PC species in the vestibular organ and cochlea. PC(16:0/20:4) was observed mainly in the vestibular sensory cells and PC(16:0/18:1) was highly localized in the nerve fiber under sensory cells. PC(16:0/16:1) was found primarily in the organ of Corti.

Conclusion
These distributional differences may be associated with the cellular architecture of these inner ear regions.

Assessment Of Methods For The Preservation Of Inner Ear Tissue By High Pressure Freezing
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Background
Electron tomography can provide detailed three-dimensional reconstructions of cellular structures at nanoscale resolution. To make the most effective use of this technique, sample preparation must preserve structures in as ‘close to life’ a state as possible with minimal artefact formation. Cryo-preparation techniques such as high pressure freezing (HPF) aim to preserve samples by freezing them rapidly at high pressure, promoting formation of vitreous water and minimising ice crystal formation. However artefacts from freezing and tissue preparation can often still be seen in these samples. Previous work has suggested that fragile tissues, such as the mammalian organ of Corti, may benefit from prefixation before HPF to improve tissue preservation for tomographic reconstruction. We have explored conditions necessary to preserve inner ear tissue for high resolution structural analysis.

Methods
For HPF, tissue was obtained from mice, gerbils and guinea pigs and mounted in aluminium planchettes with one of a variety of cryoprotectants. Some samples were chemically fixed in glutaraldehyde before HPF. Frozen samples were freeze substituted in acetone with secondary fixation and staining by osmium tetroxide, uranyl acetate and tannic acid. Warming to ambient temperature took place over 36 hours before embedding in plastic. Standard thin sections (80nm) were examined in an electron microscope operating at 80kV. Thicker sections (200nm) for electron tomography were examined in a microscope operating at 200kV. Images were collected using SerialEM software. Tomograms were reconstructed and visualised using the IMOD suite of programmes.

Results
Multiple freezing artefacts were observed in both organ of Corti and utricles that had been prepared by HPF and freeze substitution. Optimisation of cryoprotectants showed that hexadecene reduced freezing artefacts, but that no change in cryoprotectant improved the disordered actin in stereocilia that was observed. Prefixation with glutaraldehyde dramatically improved the preservation of parallel actin in stereocilia in samples frozen with hexadecene. Further improvement in preservation was observed when prefixed samples were frozen with a yeast paste cryoprotectant.
Electron tomography of samples frozen under optimised conditions revealed structural details of the interstereociliary links in a sample of guinea pig organ of Corti.

Conclusion
A chemical fixation step before HPF was found to preserve the ordered structure of actin filaments in the stereocilia far better than freezing alone. Combined with optimisation of the cryoprotectant used, this method produced samples with little evidence of freezing damage and excellent structural preservation. Electron tomography of the HPF frozen samples revealed high resolution structural details of the stereocilia bundle.

L-N-Acetylcysteine: a promising antioxidant for prevention of TNFα ototoxicity in vitro
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Background
Tumor necrosis factor α (TNFα) has been suggested to play a significant role in hearing loss following: acoustic trauma; vibration-trauma; bacterial meningitis; cisplatin exposure; and autoimmune mediated sensorineural hearing loss. Our previous work demonstrated oxidative stress in TNFα-challenged organ of Corti (OC) explants. L-N-acetylcysteine (L-NAC), an acetylated variant of L-cysteine and an excellent source of sulfhydryl (SH) groups, is converted into metabolites capable of stimulating glutathione (GSH) synthesis and acting directly as a free radical scavenger. Earlier studies have shown L-NAC to be an effective protectant against noise-induced hearing loss. The present study tests the antioxidant capability of L-NAC for protecting against TNFα-induced oxidative stress and hair cell (HC) loss.

Methods
OC explants were dissected from P-3 rats and placed in serum-free media. Explants were divided into five groups: 1) untreated controls; 2) TNFα; (2µg/ml); 3) TNFα;+L-NAC (5mM); 4) TNFα;+L-NAC (10mM); and 5) L-NAC (10mM). All explants were stained for HCs after 96hrs in vitro. After determining with HC counts that 5mM was the most effective concentration, four groups of explants were used for additional studies: 1) untreated controls; 2) TNFα; (2µg/ml); 3) TNFα;+L-NAC (5mM) and 4) L-NAC (5mM) only. Levels of: total reactive oxygen species (ROS) with CellROX®; superoxide dismutase; catalase; 4-Hydroxy-2-noneal (HNE);and total glutathione were studied after 24hrs in vitro.

Results
We observed protection of HCs from the ototoxic effects of TNFα in the L-NAC treated explants. There was no evidence of either ROS (CellROX®) or HNE immunolabeling in TNFα + L-NAC (5mM) explants compared to TNFα-challenged explants. Superoxide dismutase and catalase activities were both significantly reduced in TNFα-challenged explants compared to levels for these antioxidant enzymes present in control explants. Total glutathione levels were low in TNFα-challenged explants compared to levels for this antioxidant molecule measured in control and TNFα + L-NAC treated groups of explants.

Conclusion
Treatment of OC explants with L-NAC significantly decreases the TNFα-initiated level of oxidative stress and protected the HCs against TNFα-initiated loss. L-NAC is a promising treatment for protection of the auditory HCs undergoing inflammation-induced oxidative stress.
Action of substance P on the recovery from the intense noise exposure
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Background
Substance P is a polypeptide composed of 11 amino acids, and it is known as a perception neurotransmitter. We have already reported that the VOR gain increased after the administration of substance P into the inner ear. In addition, the nystagmus after the vestibular disorder was suppressed. In the last meeting, we have reported the role of substance P on the recovery from the temporary threshold shift after the intense noise exposure. And we have also reported that substance P affected synaptic ribbons of inner hair cells. In the present study, we administered substance P to the inner ear after TTS, and evaluated the synaptic ribbon beneath the inner hair cells.

Methods
Hartley male guinea pigs (350 - 400 g) with the normal tympanic membranes and the normal Preyer's reflexes were used in this study. Animals were divided into 2 groups (TTS group, Substance P+TTS group). After the post-auricular incision, gelatin sponge which included substance P (10-2 M) (Substance P + TTS group) or saline (TTS group) was placed on round window membrane. The mastoid bulla was closed with the dental cement. The ABR threshold was measured immediately after this operation to confirm that the hearing function did not change. The animals were exposed to the intense band noise (110 dBSPL for three hours as the TTS model). 3 hours, 12 hours, 24 hours, 3 days, 7 days later, the temporal bone was removed to evaluate the synaptic ribbon count. To ravel the synaptic ribbon, the immunohistochemistry was performed using the anti-CtBP2 antibody. These samples were observed with the fluorescence microscope.

Results
Three hours, 12 hours later, in the Substance P+TTS group, the significantly higher density of signals for synaptic ribbon were observed than those in TTS group. Twenty-four hours later, in the Substance P+TTS group, the lower density of signals for synaptic ribbon were observed than those in TTS group. There is no significant difference in the count number of a synaptic ribbon in the two groups.

Conclusion
The previous reports assumed that substance P has amplified the cochlea nerve compound action potential and that substance P protected the spiral ganglion. In the present study, we showed that the administration of substance P inhibited the change of synaptic ribbon after the intense noise exposure. The result suggests that the polypeptide promotes the recovery of the cochlear function after noise trauma.
Screening for protective effect in supplement drugs using the zebrafish lateral line
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1Yamaguchi University Graduate School of Medicine

Background
The zebrafish lateral line is a powerful system for studying hair cells and hair cell death. Hair cells can be easily labeled and imaged in vivo with fluorescence microscopy. We have previously described a screening system to rapidly assess drugs for possible ototoxic effects in anti-cancer drugs (Hirose et al., 2011). Also it is possible to screen protective effect against some ototoxicity drugs. There are many kind of supplement in USA, Europe, and Japan, etc. Supplement can be a prophylactic treatment especially against age-related hearing loss. Some countries have trouble against medical expenses, so it can be useful to such countries.

Methods
We have screened the supplement drugs for protective effects against aminoglycoside. 5-7 dpf Zebrafish (Danio rerio) embryos of the AB wild type strain were used in this study. Zebrafish larvae were exposed to supplement drugs (0, 1, 10, 100, 1000 uM) for 1 and 24 hours before 100 uM Neomycine for 1 hour. After that, they were fixed in 4% paraformaldehyde, incubated with anti-parvalbumin, and hair cell damage was assessed by fluorescent microscope. We made dose-response carve to evaluate protective effect against Neomycine. Then we confirm whether screened drug can protect inner ear hair cell on noise-induced hearing loss in guinea pigs. The ABR threshold was examined under anesthesia. The sound stimuli consisted of 2-, 4- and 8-kHz tone bursts. ABR threshold was defined as the lowest stimulus intensity to produce a liable waveform of 3-5 peaks.

Results
Dose-response carve show some of drugs show (e.g. Quercetin) protective effect against neomycin. In the guinea pigs study, we observed that the ABR threshold shift was significantly less in the Quercetin group than in control group. In addition, missing outer hair cells was lower in the Quercetin group than in the control group.

Conclusion
We show that some of supplement drugs show protective effect against neomycin, further more, screened drugs by zebrafish lateral line show protective effect against sound exposure. It is useful to screen drugs on zebrafish hair cell damage model, because zebrafish hair cell is similar not only to morphologically but also biochemically.

Cholesterol Homeostasis and Auditory Function
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Background
Animal models and human studies support a link between altered cholesterol homeostasis and sensorineural hearing loss, implicating pathologies that affect the cochlea, auditory nerve and/or central auditory pathways. However, knowledge of the relative quantities and species of lipids in the cochlea, as well as how circulating lipids modulate cochlear cellular activities, is currently lacking. The rising prevalence of obesity and hypercholesterolemia demands a better understanding of how systemic cholesterol homeostasis influences cochlear cholesterol homeostasis and hearing function. We hypothesize that cholesterol homeostasis is essential for hearing function and maintenance, and that changes in cholesterol levels lead to decreased cochlear and hearing function.

Methods
Auditory function was assessed by measuring auditory brainstem responses (ABRs) as described previously by our group. Total cholesterol levels in cochlear tissues were quantified using the Amplex Red cholesterol assay (InVitrogen, Carlsbad, CA) and compared to serum lipid panels. Stria vascularis capillary organization and perivascular-resident macrophage-like melanocytes (PVM/Ms) were visualized by whole-mount staining and fluorescence confocal microscopy.

Results
We have demonstrated that cochlear levels of total cholesterol and serum lipid panels change significantly during normal maturation of cochlear and auditory function. Interestingly, we have shown that the sensory epithelium (SE) contains higher levels of cholesterol than the vascular compartment. Current studies focus on how systemic hypercholesterolemia affects the hearing system by using the Low-Density Lipoprotein Receptor (LDLR) KO mouse model. LDLR KO mice display elevated hearing thresholds, indicative of hearing loss. In addition, LDLR KO mice have altered organization of the vasculature within the stria vascularis, the metabolic engine of the cochlea. We are also exploring how these changes
develop with age, as well as the contribution of PVM/Ms to maintenance of the structural integrity of the blood-labyrinth-barrier (BLB) in the stria vascularis capillaries.

**Conclusion**

Preliminary results not only suggest that serum cholesterol levels influence cochlear cholesterol homeostasis, but also the existence of a mechanism for concentrating cholesterol in the SE or endogenous synthesis of cholesterol by hair cells, supporting cells or other cells of the organ of Corti. Hypercholesterolemic LDLR KO mice have impaired auditory function, which may result from altered stria vascularis structure or altered cochlear cholesterol homeostasis.

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**Activation of Endoplasmic reticulum stress in an inner-ear immortomouse cell line**

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**Background**

Induction of mild endoplasmic reticulum (ER) stress has been associated with an increased tolerance to cellular toxins. But under severe ER stress, specific stress-associated apoptotic pathways are activated. We investigated the alteration of HEI-OC1 cell under activation of ER stress.

**Methods**

ER stress was induced after exposure to tunicamycin and thapsigargin. We examined the unfolded protein response (UPR) by measuring the protein levels of specific genes such as glucose-regulated protein of 78kDa (GRP78) and GRP94 in HEI-OC1 cell.

**Results**

ER stress inducers resulted in a dose-dependent increase of cell death, but the magnitude of cytotoxicity was inducer-dependent. Cytotoxicity was not increased by mild ER stress. GRP78 and GRP98 expression were statistically increased after activation of ER stress.

**Conclusion**

Our results indicate UPR and cytotoxicity are dependent on the ER stress inducer in HEI-OC1 cell. These results suggest that the ER stress may play diverse roles in ototoxicity of HEI-OC1 cell.

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**Evaluation of the ototoxicity of Gentian violet solution**

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**Background**

Antibiotic abuse can lead to an increase in the incidence of multiple antibiotic resistance among organisms. Non-antibiotic solutions such as Burow's solution and Gentian violet solution have been reported to be effective in treating refractory otorrhea caused by multiple drug-resistant bacteria or fungi. Burow's solution has astringent and antibacterial properties while Gentian violet solution has a strong sterilizing effects. Recent studies have demonstrated that intratympanic application of Burow's solution causes degeneration of the outer epithelium and outer hair cell loss in the basal turn in guinea pigs. Although Gentian violet solution is also known to show ototoxicity, there is very little data regarding the physiological and morphological changes following intratympanic application of Gentian violet solution. We aimed to examine the effects of Gentian violet solution.

**Methods**

Albino guinea pigs with normal Preyer's reflex were used. All animals were anesthetized with an intramuscular injection of xylazine and ketamine. ABR and DPOAE were recorded at 4, 8, 16,32 kHz. Under aseptic conditions, an incision was made in the left postauricular region, and a small hole was made in the tympanic bulla to allow direct visualization of the RWM. In control animals, 0.05 ml of physiological saline was directly dropped onto the RWM. In experimental animals, 0.05 ml of 1% Gentian violet solution was directly dropped onto the RWM. Seven days later, ABR and DPOAE were recorded, and the left temporal bone was removed. In half of the experimental animals, the cochleae were dissected using the surface preparation technique, stained with rhodamine-phalloidin, and examined under a fluorescence microscope. In
the remaining experimental animals, the cochleae were embedded in paraffin and section in the radial plane at 4-7µm using a microtome.

**Results**

In the control animals, no marked ABR threshold shifts were observed at any frequency. No hair cell loss was observed anywhere in the organ of Corti. In the experimental animals, the ABR thresholds shifts at all frequencies were significantly higher than those in the controls. On the other hand, DPOAE showed normal responses at all frequencies. Many phalangeal scars were observed in the region of outer hair cells in the lower half of the basal turn.

**Conclusion**

Gentian violet solution applied onto RWM may cause marked hearing impairment. This hearing impairment may be induced not only by outer hair cell loss but also by damage to the spiral bundle and/or spiral ganglion.

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**CCR2 expression defines cochlear response to LPS preconditioning**

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**Background**

Lipopolysaccharide (LPS) at low doses has been widely used as a preconditioning stimulus to prime the immune system before a subsequent injury. In ototoxicity, LPS pretreatment worsens hair cell loss and exacerbates threshold shift caused by kanamycin/furosemide combination therapy. This LPS induced exacerbation of hearing loss is reversed, and LPS pretreatment is protective against threshold shift in mice that are deficient for CC-chemokine receptor 2, CCR2, which is expressed in inflammatory monocytes that enter the inner ear after exposure to LPS.

**Methods**

8 weeks old C57Bl6 mice were injected with saline or LPS (0.5mg/kg/day for two consecutive days) and then injected with kanamycin/furosemide (1000mg/kg and 180 mg/kg respectively) on the third day. ABR thresholds were obtained at day five after kanamycin/furosemide, and cochleas were harvested and analyzed for hair cell survival and numbers of monocyte/macrophage entry and chemokine receptor expression.

**Results**

Threshold shift after LPS priming is about 40 dB greater than threshold shift in mice injected with saline prior ototoxic injury in CCR2+/+ mice. CCR2 expressing monocytes are recruited in abundance to the cochlea with LPS pretreatment, but few CCR2 expressing monocytes are present in the inner ear after ototoxic exposure alone. Threshold shift in LPS-pretreated mice was about 20 dB lower compared to saline-pretreated mice exposed to the same kanamycin/furosemide treatment when CCR2/-/- mice were studied. Thus, LPS in CCR2 wild-type mice has the effect of exacerbating ototoxicity, while in CCR2 null mice, LPS is protective.

**Conclusion**

CCR2 expression in peripheral monocytes is critical in defining whether LPS pretreatment causes exacerbation of ototoxic injury or mediates protection. This finding represents an important example of where mononuclear phagocytes, recruited into the inner ear can play an important role in determining hearing function after hair cell injury.
Selective Hair Cell Ablation Recruits Macrophages into the Cochlea and Activates Microglia in the Cochlear Nucleus

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Background
Prior studies have demonstrated that macrophages are recruited into the cochlea after acoustic trauma or aminoglycoside ototoxicity. However, the signals that initiate macrophage recruitment, as well as the contribution of macrophages to cochlear pathology and/or repair, have not been determined. Injury to the cochlea also activates microglia within the cochlear nucleus, but the role of those cells in brainstem pathology or plasticity is also unclear. We have used a novel mouse model to characterize the response of both cochlear macrophages and microglia in the cochlear nucleus to selective elimination of hair cells.

Methods
Mice expressing diphtheria toxin receptor under control of the Pou4f3 promoter were crossed to CX3CR1-GFP mice, which express GFP in all macrophages and microglia. Mice 6-8 weeks of age were given single IM injection of diphtheria toxin (DTX, 25ng/20gm), which caused an extensive loss of both inner and outer hair cells. At 1-12 days after DTX treatment, mice were euthanized and perfused with 4% paraformaldehyde. Cochleae and brainstems were isolated and processed for immunohistochemical labeling of neurons and hair cells.

Results
Hair cell loss was first evident at 3 days after DTX injections. At 7 and 12 days post-injection, we observed nearly complete ablation of both inner and outer hair cells. This was accompanied by recruitment of numerous GFP-expressing macrophages into the cochlea. Notably, such macrophages were abundant in the scala tympani, immediately below the basilar membrane. Those cells often extended processes into the injured outer hair cell region. Macrophages are also observed in close contact with dendrites of the spiral ganglion neurons within the habenula perforata and with neurons crossing the tunnel of Corti and extending towards outer hair cells. Also, macrophages in the DTX injected mice are amoeboid compared to the ramified cells in DTX injected wild-type mice. We also observed many GFP-expressing microglia within the cochlear nucleus of mice status post hair cell ablation. Increased numbers of microglia were present in the ventral cochlear nucleus at 12 days after DTX injection. Finally, injection of diphtheria toxin into wild type mice did not induce an inflammatory response in the cochlea or auditory brainstem.

Conclusion
The mouse model employed in these studies permits the selective complete elimination of hair cells, without any other evident pathology. Our data indicate that the loss of cochlear hair cells is sufficient to recruit macrophages into the cochlea and microglial activation within the cochlear nucleus.

Inflammation mobilizes vestibular perivascular resident macrophage-like melanocytes

Fei Zhang¹, Lingling Neng¹, Allan Kachelmeier¹, Xiaorui Shi¹
¹Oregon Hearing Research Center

Background
A large number of perivascular cells expressing both macrophage and melanocyte characteristics (named perivascular resident macrophage-like melanocytes, PVM/M) were previously found in the intra-strial fluid-blood barrier.

Methods
In this study, we used confocal microscopy combined with fluorescence immunohistochemistry approach.

Results
The PVM/Ms were found to also locate in the area of the blood-labyrinth barrier of the vestibular system in normal adult cochlea, including in the three ampullae of the semicircular canals (posterior, superior and horizontal), utricle, and saccule. The cells were identified as PVM/Ms, as they were positive for the macrophage and melanocyte marker proteins F4/80 and MIFT. Similar to the PVM/Ms present in the stria vascularis, the PVM/Ms in the vestibular system are closely associated with microvessels and structurally intertwined with endothelial cells and pericytes. The density of PVM/Ms in normal, unstimulated tissue is 225 ± 43/mm² in the utricle, 191 ± 25/mm² in the saccule, 212 ± 36/mm² in horizontal ampullae, 238±36/mm² in anterior ampullae, and 223±64 /mm² in posterior ampullae. Injection of bacterial lipopolysaccharide (LPS, 5 mg/ml) into the middle ear through the tympanic membrane dramatically increases the presence of PVM/Ms irregularly situated along capillary walls in all regions within a 48 hour period. PVM/M foot
processes were notably detached from some of the capillaries. The inflammatory response significantly increased vascular permeability and leakage.

**Conclusion**
The results indicate PVM/Ms in the vascular wall of the blood-labyrinth barrier are contributing factors in the local inflammatory response. This work was supported by National Institutes of Health grants NIH NIDCD R01-DC010844 (XS), DC R21DC1239801 (XS), and NIHP30-DC005983.

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**Pigment epithelium-derived growth factor protects against noise-induced hearing loss**
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**Background**
Acoustic trauma not only directly damages sensory hair cells, but it also disrupts the cochlear intra-strial fluid-blood barrier in the stria vascularis. An intact intra-strial fluid-blood barrier is critical for sustaining the ionic gradients and endolymphatic potential (EP) essential for hearing function.

**Methods**
In this study, animals were exposed to broadband noise at 1170 dB SPL in a sound exposure booth for three hours and for an additional three hours the following day. An auditory brain-stem response audiometry to pure tones was used to evaluate hearing function in control, noise-exposed, noise-exposed + pigment epithelium-derived growth factor (PEDF), a potent endogenous inhibitor of vasopermeability, treated groups. Endocochlear potential (EP) was recorded. Expression of tight junctions were detected with immunohistochemistry combined with confocal microscopy.

**Results**
We found that PEDF protects against noise-caused hearing loss. Animals pre-treated with PEDF show substantially reduced EP drop with noise exposure. The PEDF promotes expression of tight junction-associated proteins such as ZO-1 and VE-cadherin between endothelial cells and stabilizes accessory perivascular-resident macrophage-like melanocytes by increasing expression of neural cell adhesion molecule (NCAM, CD56).

**Conclusion**
Our results demonstrate the critical role of PEDF in maintaining cochlear homeostasis and suggest its potential therapeutic role in protecting against noise-induced vascular permeability and hearing loss. This work was supported by National Institutes of Health grants NIH NIDCD R01-DC010844 (XS), DC R21DC1239801 (XS), and NIHP30-DC005983.
IL-10 Inhibits Otitis Media-induced Cochlear Inflammation via HMOX1-dependent NF-κB Inactivation

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Background

Recently, we have demonstrated that the spiral ligament fibrocytes (SLFs) play a pivotal role in the cochlear inflammation secondary to otitis media (OM). However, considering that the middle ear infection is one of the most common infectious diseases in children, clinical evidence of the cochlear symptoms secondary to OM is less than our expectation. This suggests that cochlear inflammation is tightly regulated to prevent tissue damage by excessive inflammatory response. In this study, we aim to determine the molecular mechanism involved in IL-10-mediated inhibition of OM-induced cochlear inflammation.

Methods

To localize IL-10-expressing cells, immunolabeling and RT-PCR analysis were conducted. To determine MCP-1 regulation, we performed qRT-PCR analysis and luciferase assays. HMOX1 was activated by CoPP and CORM and was inhibited using the specific siRNA. Chromatin immunoprecipitation (ChIP) analysis was performed to determine binding of NF-κB to the enhancer of the MCP-1 gene.

Results

The SLFs appeared not to produce IL-10, but we found that IL-10-expressing cells are localized in the spiral ligament. The SLFs were shown to inhibit NTHi-induced MCP-1 expression upon exposure to IL-10, which was suppressed by silencing of IL-10RA. The inhibitory effect of IL-10 was mimicked by treatment of CoPP and CORM but was blocked by silencing of HMOX1. The SLFs were found to activate NRF2 in response to IL-10, resulting in up-regulation of HMOX1. Luciferase assays showed that IL-10 inhibits NTHi-induced NF-κB activation. Furthermore, ChIP analysis demonstrated that IL-10 inhibits NTHi-induced binding of p65 to the NF-κB binding motif of the MCP-1 gene.

Conclusion

Taken together, our results suggest that IL-10 modulates OM-induced cochlear inflammation through HMOX1-mediated negative regulation of MCP-1 expression in the SLFs. [Supported by NIH grants: DC005025, DC006276 and DC011862]
Interaction of IFN-gamma and TNF-alpha in Cisplatin Ototoxicity
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Background
Inflammatory response is known to be involved in cisplatin cytotoxicity. Previously, we have demonstrated that the auditory hair cells induce pro-inflammatory cytokines in response to cisplatin. Moreover, inhibition of cytokines was shown to ameliorate cisplatin ototoxicity. However, we poorly understand interactions of cytokines in cisplatin ototoxicity. In this study, we aim to determine a role of IFN-γ and TNF-α in cisplatin ototoxicity.

Methods
To determine cell viability, we performed MTT assays and cytometric analysis with trypan blue staining. For the ex vivo model of ototoxicity, the organ of Corti was isolated from the Math-1-EGFP mice and explanted on the PCTE membrane. Superoxide generation was monitored using flow cytometric analysis with hydroethidine staining. qRT-PCR analysis was conducted to determine NOX1 regulation.

Results
TNF-α appeared to augment cisplatin-induced ototoxicity in vitro and ex vivo, which was blocked by a small molecule inhibitor of TNF-α activity. IFN-γ alone appeared to insignificantly enhance cisplatin-induced cytotoxicity in the HEI-OC1 cells. However, pretreatment of IFN-γ was found to highly potentiate TNF-α-mediated enhancement of cisplatin cytotoxicity. In addition, IFN-γ-mediated enhancement of TNF-α-induced cytotoxicity was inhibited by Epigallocatechin-3-gallate, indicating the involvement of STAT1 signaling. The HEI-OC1 cells appeared to up-regulate superoxide generation and NOX1 expression in response to TNF-α, which was enhanced by the pretreatment of IFN-γ. Cisplatin-induced caspase-3 activation was enhanced upon exposure to TNF-α in the HEI-OC1 cells. Pharmacological inhibition of caspases was found to ameliorate TNF-α-induced enhancement of cisplatin cytotoxicity.

Conclusion
Taken together, our results suggest that interaction of IFN-γ and TNF-α augments cisplatin ototoxicity, which involves superoxide generation. [Supported by NIH grants: DC005025, DC006276 and DC011862]
Rescue of Hearing and Vestibular Defects in Usher Syndrome Using Antisense Oligonucleotides: Histological and Scanning EM Analysis

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LSU HSC, Rosalind Franklin Chicago Medical School, Isis Pharmaceuticals

Background

Usher syndrome (Usher) is the most common cause of hereditary deaf-blindness, characterized by sensorineural hearing impairment combined with retinitis pigmentosa, and in some cases, vestibular dysfunction. The only treatment for Usher is the use of prosthetics, such as hearing aids and cochlear implants, and the feasibility of curing deafness at the cellular level is uncertain. A mouse model for Usher has been developed based on the Acadian Usher mutation, in which the USH1C.216G>A mutation (216A) has been knocked into the mouse Ush1c gene. The 216AA mice have profound deafness, vestibular dysfunction and develop retinal degeneration characteristic of Usher patients. The 216A mutation introduces a cryptic splice site that is used preferentially over the authentic site producing a truncated mRNA and protein product. We hypothesize that blocking the mutant splice site with antisense oligonucleotides (ASOs) will activate the authentic site, rescue protein expression and be therapeutic for mice and ultimately Usher patients.

Methods

ASO activity was optimized using a cell-free splicing system, an ASO-tiling screen and cell lines isolated from the 216AA mice and an Usher patient with the 216AA mutation. 216AA mice were injected with ASOs and correction of splicing and protein expression was quantitated by rt-pcr and western blot, respectively. Harmonin b expression and localization was examined by immunohistochemistry. Outer hair cell (OHC) bundle morphology was evaluated by scanning electron microscopy. Hearing was evaluated by auditory-evoked brainstem response (ABR) analysis.

Results

Treatment to neonatal 216AA mice with a single systemic dose of ASOs partially corrects splicing and protein expression in the cochlea, improves OHC stereocilia organization and rescues cochlear hair cells, vestibular function and hearing function.

Conclusion

Our results demonstrate the therapeutic potential of ASOs in Usher syndrome and other diseases caused by mutations that disrupt splicing.
Trafficking Deficiency of KCNQ4 Channels Associated with Progressive Sensorineural Hearing Loss Can be Corrected by Molecular Chaperone HSP90β
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Background
The inner ear KCNQ4 channel plays crucial roles in maintaining cochlear ion homeostasis and regulating hair cell membrane potential that are essential for normal auditory functions. Dysfunction of the KCNQ4 channel leads to the autosomal dominant non-syndromic deafness DFNA2, characterized by progressive sensorineural hearing loss. At younger ages, hearing loss in DFNA2 patients is moderate and predominantly at high frequencies. At older age, however, all affected individuals have severe to profound hearing impairment with all frequencies involved. Genetic studies have identified numerous KCNQ4 mutations in DFNA2 patients. However, the molecular etiology of DFNA2 is elusive.

Methods
Using immunofluorescent, biochemical, and electrophysiological approaches, we investigated functional consequences of seven DFNA2 mutations.

Results
Dramatic decrease in cell surface expression of KCNQ4 channels was detected by immunofluorescent microscopy and confirmed by Western blot for all mutations tested, while the cellular levels of these mutant channels remained normal. In addition, none of these mutations interrupt subunit interaction, consistent with their dominant-negative effects on the wild type KCNQ4 trafficking. Moreover, we found that upregulation of HSP90β, a major molecular chaperone that controls the KCNQ4 biosynthesis, significantly improved cell surface expression of DFNA2 mutants; KCNQ4 expression in the plasma membrane was restored in cells mimicking heterozygous conditions of DFNA2 patients. However, no significant changes in KCNQ4 currents was observed after restoration of KCNQ4 surface expression.

Conclusion
Trafficking deficiency and impaired KCNQ channel function are two underlying mechanisms for hearing loss in DFNA2.
Oral Capsaicin Consumption Protected Against Noise Induced Hearing Loss (NIHL)
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Background
Exposure to high levels of noise is the most common cause of hearing loss in adults. Noise exposure leads to cellular stress in the cochlea by activating and increasing a cochlear specific NADPH oxidase, NOX3 and transient receptor potential vanilloid 1 (TRPV1) channel. Increased NOX3 and TRPV1 expression lead to increased inflammation as evidenced by increases in proinflammatory mediators such as tumor necrosis factor-α (TNF-α), inducible nitric oxide synthase (iNOS) and cyclooxygenase-2 (COX-2). This mechanism of reactive oxygen species (ROS)-induced inflammation leading to hearing loss is similar to that underlying cisplatin ototoxicity, previously delineated in our laboratory. We have previously shown that direct activation of TRPV1 by capsaicin reduced cisplatin ototoxicity in the rat. Therefore, in this study, we assessed the effectiveness of capsaicin against noise-induced hearing loss (NIHL).

Methods
Baseline auditory brainstem responses (ABRs) of male Wistar rats were recorded. This was followed by intra-tympanic (IT) (50µl of 0.1µM capsaicin) or the oral (0.01 to 2% in water) capsaicin administration and noise exposures (122dB octave band noise centered at 16kHz for 1h). Post-treatment ABRs were recorded 21 days following noise exposures. Rat cochleae were extracted and used for various assays. Functional assessments include ABRs and scanning electron microscopy (SEM). Gene expression studies were conducted using real time real time PCR and by immunohistochemistry (IHC) of the mid-modiolar sections of the rat cochleae.

Results
Our data indicate that noise exposure produces a 45-60dB shift in ABR thresholds. Rats administered oral capsaicin demonstrated significantly reduced ABR threshold shifts of <10dB. Significant decreases in ABR threshold shifts following noise exposures were also observed following IT capsaicin to <30 dB. Capsaicin-mediated otoprotection was co-related with preservation of outer hair cells, decreased expression of NOX3 and TRPV1 genes and inflammatory mediators such as TNF-α, iNOS and COX-2.

Conclusion
We conclude that noise trauma increases cochlear ROS generation and pro-inflammatory genes in the cochlea. In addition, pretreatment with capsaicin (either orally or IT) reduced the responses to noise trauma and protected against NIHL. Our study suggests that the consumption of capsaicin in spicy food could offer protection against NIHL.
**Oral EpigallocatechinGallate (EGCG) Protects Against Cisplatin-Induced Ototoxicity in Rats**

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**Background**

Cisplatin is an anticancer drug used to treat various solid tumors. Patients treated with cisplatin develop significant ototoxicity and nephrotoxicity. Various drugs are under investigation for the alleviation of cisplatin-induced ototoxicity and nephrotoxicity. In previous studies, we reported that the cisplatin triggers reactive oxygen species (ROS) generation, activation of signal transducer and activator of transcription-1 (STAT-1) and inflammation in the cochlea, which ultimately leads to damage and death of outer hair cells (OHCs) and hearing loss. Accordingly, inhibition of STAT-1 by siRNA or inflammation by tumor necrosis factor (TNF-α) antagonist, etanercept, protected against death of OHCs and preserved hearing. In this study, we test the ability of the green tea extract, epigallocatechingallate (EGCG), a known inhibitor of STAT1, to protect against cisplatin-induced cell death in vitro and against hearing loss in Wistar rats.

**Methods**

Organ of Corti-derived cells (UB/OC-1) were cultured and treated with cisplatin in the absence and presence of EGCG to determine effects on STAT1 activity and cell apoptosis. Male Wister rats (200-250g) were treated with oral EGCG (100mg/kg), followed by cisplatin (11mg/kg, i.p.). Auditory brainstem responses (ABRs) and scanning electron microscopy (SEM) studies were performed to assess hearing loss and OHC morphology. University of Michigan squamous cell carcinoma (UMSCC) 10b cells were used to test for potential antitumor interference by EGCG.

**Results**

Western blot studies indicated that EGCG inhibited cisplatin-induced STAT1 phosphorylation and activation. EGCG also inhibited cisplatin-induced p53 expression and increased the levels of anti-apoptotic proteins, Bcl-2 and Bcl-xl. In vivo studies indicated that EGCG protected against cisplatin-induced loss of OHCs. Current studies are testing the effects of EGCG against cisplatin-induced loss of outer hair cells. Studies in UMSCC 10b cells demonstrated that EGCG augmented cisplatin-induced cell killing, indicating a potential lack of anti-tumor interference.

**Conclusion**

We propose that EGCG could serve as an important treatment for cisplatin-induced hearing loss by inhibiting a STAT-1 dependent apoptotic pathway. Furthermore, EGCG did not interfere with cisplatin-induced killing of squamous cell carcinoma cells. These studies would support the use of oral EGCG for the treatment of cisplatin ototoxicity. (Supported by NIH grants R01Dco2396 and R15DC011412 and funds from SIU SOM).

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**Effect of Non-steroidal Anti-Inflammatory Drugs (NSAIDs) on Prestin.**

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**Background**

High doses of salicylate can cause reversible hearing loss and tinnitus. According a classification of ototoxicity-inducing drugs by Cianfrone and colleagues, other NSAIDs can trigger side effects related with hearing and cause tinnitus. Although ototoxicity can be due to the interaction of these medicines with any sensitive part of the hearing system, we are investigating a possible prestin-related mechanism for these adverse reactions.

**Methods**

To measure the non linear capacitance of prestin, we used HEK-cells stably transfected with a plasmid containing the prestin gene. The NLC was measured from these cells after 48 hours of induction. For each cell, a control NLC in the absence of drug was obtained. The appropriate NSAID was then perfused in the recording chamber and a second NLC was measured. We also investigated the impact of NSAIDs when present only outside or only inside the cell to address a possible sidedness of the effect.

**Results**

We assessed the effect on prestin of the widely used ibuprofen, acetaminophen and naproxen, all of which have ototoxic side effects. We also report the impact of salicylate-derivatives diflusinal and piroxicam whose side effects on hearing are stronger than with most NSAIDs. All NSAIDs tested, except piroxicam, triggered a voltage shift of the NLC. The stronger shift in was observed with diflusinal, which triggered a 46mV shift at 2 mM. Moreover, a change in charge density was observed when ibuprofen or diflusinal were perfused. The presence of 2 mM diflusinal decreased the charge density by
over 50% which could suggest a competition with Cl- ions. The perfusion of 6mM ibuprofen, on the other hand, increased the charge density by 20%, suggesting a decrease in membrane lateral pressure.

Conclusion
NSAIDs induce alterations in the function of prestin which could impact hearing. The NSAID-induced shift in the NLC could modify the function of the OHC at resting potential and alter cochlear amplification. We also witnessed a correlation between the $V_{1/2}$-shift and the pKas of the NSAIDs. The drug that shows no effect was the least charged (piroxicam), the ones that trigger hyperpolarization were positively charged (ibuprofen, naproxen and diflusinal) while the negatively charged molecule (acetaminophen) had a depolarizing effect. This suggests that the charges brought to the membrane by the NSAIDs are responsible for the alteration in $V_{1/2}$.

Mechanical Tuning in the Hearing System of Bushcrickets
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Background
The high-frequency hearing organ (crista acustica, CA) of bushcrickets is located within the tibiae of the forelegs. Earlier studies revealed a tonotopical representation of sound-induced amplitude maxima along the organ when stimulated with 80 dB SPL. The anatomical basis for tonotopy, however, is unknown. It has been assumed that the varying mass of supporting cells on top of each dendrite, called cap cells, might influence sensory cell tuning by acting as mechanical resonators. To investigate the influence of possible mechanical resonances, we analyzed the sound-induced mechanical tuning of the CA as well as the anatomy of the hearing system.

Methods
The tropical bushcricket Mecopoda elongata was used to investigate mechanical vibration thresholds. For this purpose, pure-tone stimuli (2-77 kHz, 10-80 dB SPL) were applied and the mechanical vibration response of the CA was measured by a laser-Doppler-vibrometer. Additionally, we examined the anatomical features of the hearing system by analyzing the hearing organ structures on cross sections through the CA.

Results
In a first series of experiments the mechanical response to a set of different frequencies and levels was determined at three different locations along the hearing organ (proximal, medial and distal). According to the predicted tonotopical preference of the sensory cells, low frequencies elicited the most sensitive response in the proximal region and high frequencies in the distal region. The tuning curves from the proximal and medial locations revealed distinct characteristic frequencies at stimulus frequencies of 7 and 17 kHz, respectively. In contrast, in the distal region no distinct characteristic frequency could be identified. Anatomical investigations of the CA revealed a steady decrease in size of all relevant hearing-organ structures from the proximal to the distal end supporting the tonotopical distribution of displacement amplitude maxima.

Conclusion
Mechanical threshold curves measured at a proximal and medial location along the CA revealed a tonotopical tuning of sensory cells supported by the anatomical gradient along the CA. Nevertheless, the fine tuning of mechanical CA vibration, especially the broad tuning curve with no distinct characteristic frequency measured at the distal region, seems to be influenced by unknown additional factors.
Phase-Locking of Individual and Coupled Hair Bundles of the Bullfrog Sacculus
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Background
Bullfrog sacculus has been shown to detect subnanometer-level mechanical signals. This exquisite sensitivity potentially arises from the synchronization between hair bundles, mechanically coupled via the otholitic membrane under in vivo conditions. In this work, we study the phase-locking behavior of single and coupled hair bundles to external stimulations.

Methods
Calibrated glass fibers are attached either to individual hair bundles, or positioned to couple to two bundles simultaneously. The fibers are then used to deliver sinusoidal mechanical stimulation. The phase of the oscillations is extracted from the phase portrait of displacement and velocity.

Results
At low stimulus amplitudes, the phase of hair bundle oscillation exhibits diffusive behavior, with a decrease in variance with increasing stimulus level. As the stimulus becomes stronger, phase-locking can be observed with the appearance of phase slips. We observe that phase-locking occurs at ~ 10 nm stimulus, the level which corresponds to the threshold of compressive nonlinearity. In the absence of external stimulation, mechanically coupled hair bundles exhibit synchronization.

Conclusion
Individual hair bundles exhibit signatures of entrainment, evidenced by peaks in their phase histograms, at stimuli as small as ~ 2nm. With increasing stimulus amplitude, phase-locking occurs and phase slips can be observed; a behavior that is indicative of a saddle node on invariant circle (SNIC) bifurcation. We also find strong correlation between the occurrence of phase slips and the regime of compressive nonlinearity.
Statistical Properties of Spontaneous Hair Bundle Oscillations
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Background
Under in vitro conditions, hair bundles in the turtle papilla and the bullfrog sacculus have been shown to oscillate spontaneously when decoupled from the overlying membrane. This spontaneous motility results from the interaction of two mechanisms that affect the bundle’s position and which act on different time scales. Here we analyze long, multi-minute recordings of the position of freely oscillating hair bundles to extract statistics on their variation.

Methods
Spontaneously oscillating hair bundles in freshly dissected sacculi of the American bullfrog (R. catesbeiana) were filmed continuously at a high frame rate (500 fps) for several minutes. The position of a hair bundle was extracted from each image in these recordings by fitting a Gaussian distribution to its intensity profile. We developed an automated routine which examined the local variation in amplitude and its derivative to detect bundle oscillations within each trace.

Based on the detected oscillations, several of their parameters were extracted: duration, amplitude, and inter-oscillation interval. Owing to the length of the recordings, over 2000 oscillations were present in any given trace. This results in rigorous statistical characterization of the oscillation’s properties, allowing us to investigate the role of noise.

Results
Histograms of inter-oscillation intervals are either uni- or bimodal, resulting from “regular” and “bursting-type” oscillation patterns, respectively. For individual oscillations, we found that their duration increases with the quiescent interval immediately preceding them. This is most evident in bursting-type oscillations, where the first oscillation in a burst (group of closely spaced oscillations) takes longer than the rest.

The length of the recordings also permitted the calculation of their correlation dimension, which can be used to determine whether the underlying system exhibits chaos. Preliminary calculations indicate that the correlation dimension grows with the embedding dimension, M.

Conclusion
The long recordings of spontaneously oscillating hair bundles facilitate a rigorous statistical quantification of their properties. Based on an automated detection of individual oscillations we are able to identify two types of behavior for the cell: “regular” vs. “bursting” oscillations. The latter display a Poisson-like distribution in the number of oscillations per burst. The long recordings also facilitate the calculation of the trace’s correlation dimension, which is used to test for chaotic behavior in a system. The preliminary observation that this dimension grows with the embedding dimension implies that fluctuations in the bundle’s position are stochastic. Supported by NIH grant RO1 DC011380.

IHC stereocilia are excited by at least four mechanical drives that, in combination, explain formerly anomalous auditory-nerve data
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Background
The bending of inner-hair-cell (IHC) stereocilia controls cochlear output, but the mechanisms that produce this bending are poorly understood. The common conception that IHC excitation is due only to shearing between the reticular lamina (RL) and the tectorial membrane (TM) does not explain auditory-nerve (AN) data. Two key new concepts are: (1) for low-frequency sounds, the RL-TM gap varies cyclically and produces fluid flow within the gap that bends IHC stereocilia (Nowotny and Gummer 2006, PNAS 103:2120), and (2) outer hair cell (OHC) stereocilia length can change (Hakizimana et al. 2011, ARO Abstr. #355) so the RL-TM gap over OHCs is not fixed.

Methods
We hypothesize how fluid flows drive IHC stereocilia, and show that these drives explain AN data.

Results
Last year we postulated that, in addition to classic SHEAR, there are two other IHC drives: (1) TM-PUSH: For upward BM motion, the force that moves the TM compresses the RL-TM gap over OHCs causing inward radial flow past IHCs. (2) OHC-MOTILITY: Upward basilar-membrane (BM) motion causes OHC somatic contraction which tilts the RL, compresses
the RL-TM gap over IHCs and expands the RL-TM gap over OHCs, thereby producing an outward radial fluid flow. For upward BM motion these drives deflect IHC stereocilia in the inhibitory and excitatory directions, respectively. We now add a second drive that is inhibitory for upward BM motion, the CILIA-SLANT drive: Motions that produce large tilting of OHC stereocilia squeeze the supra-OHC RL-TM gap and cause inward radial flow past IHCs. Combinations of these drives explain: (1) the reversal at high sound levels of AN initial peak (ANIP) responses to clicks, and medial olivocochlear (MOC) inhibition of ANIP responses below, but not above, the ANIP reversal, (2) dips and phase reversals in AN responses to tones in cats and chinchillas, (3) hypersensitivity and phase reversals in tuning-curve tails after OHC ablation, and (4) MOC inhibition of tail-frequency AN responses.

Conclusion
The ability of these IHC drives to explain previously anomalous AN data provides strong, although indirect, evidence that these drives are significant. The OHC-MOTILITY drive is another mechanism, along with BM motion amplification, that uses active processes to enhance cochlear output. Overall, the success of the hypotheses presented here argues for a new view of how the cochlea works at frequencies below 3 kHz.

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The effect of noise-clicks on IHC bundle mechanics
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Background
Mechanotransduction, the process by which the hair cells convert mechanical stimuli into a electrical response, is managed by transduction (ion) channels located at the attachment points of the tip links. The transduction channels are opened by elastic elements, i.e. gating springs, and are stretched when the hair bundle is deflected toward its tall edge. A molecular gate that opens and closes based on the tensioning in the spring controls each channel. Noise-induced damage the hair bundles of the inner ear appears as disarray, fusion, or collapsing of the stereocilia. This damage may result from breakage of side links and tip links.

Methods
The effects of noise clicks on IHC stereocilia bundle mechanics were investigated at several frequencies in the apical turn of a guinea pig temporal bone preparation using time-resolved confocal imaging and optical flow algorithms. The experimental data was used with the Smith-Chadwick computational model to investigate the effect of the noise-clicks on stereocilia tip link tensioning and bundle mechanics.

Results
Preliminary data show that the noise clicks reduced the stereocilia bundle motion amplitude and this effect was also consistent with a decrease in the cochlear microphonics.

Conclusion
The results of this study will show that the noise-click exposure may affect mechanotransduction through disruption of stereocilia mechanics.

Prolonged deflections suppress active hair bundle motion
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Background
Active hair bundle motility is one of the signatures of the amplification mechanism in non-mammalian hair cells. Here we impose high-amplitude deflections on the hair bundle for 0.1-50 seconds. Due to incompleteness of the adaptation mechanism, this is expected to change the opening probability of the transduction channels, thus mechanically bringing the bundle out of its equilibrium state. Recovery of hair bundle motion was characterized, and the effects of calcium on the feedback mechanism were investigated.

Methods
Adult bullfrogs (RanaCatesbiana) were anesthetized and decapitated following the UCLA DLAM protocol to extract the sacculus in the inner ear. The epithelium was then placed in a two-compartment chamber to separate the solutions
bathing the apical and basal side, artificial endolymph and perilymph respectively. Mechanical stimuli were applied using two techniques. For one set of experiments, a glass probe, made with a micropipette puller, was used to deflect the bundles. In another set of measurements, magnetic beads of ~ 1 micron diameter were attached to the stereociliary bundles, and a magnetic probe was used to impose deflections. To study the influx of calcium, hair cells were dissociated after incubation of protease XXIV for 10 minutes and papain for 20 minutes, and loaded with Fluo-4 AM.

**Results**
Traces of hair bundle motility recorded immediately post stimulation show a slow drift towards the equilibrium position, without oscillatory motion. Bundles recover their innate oscillation between 0.1 – 1 second after cessation of the stimuli, and the quiescent time (Tq) shows a strong dependence on stimulus duration. Decreased calcium concentration leads to a slower spontaneous oscillation, consistent with prior results in the field. The quiescent interval induced by deflection of hair bundles in a low-calcium environment is shorter, recovering more rapidly post stimulus cessation.

**Conclusion**
High-amplitude stimulus induces a quiescent interval in hair bundles, with a slow relaxation that exhibits two characteristic time scales. The two time constants of the recovery show a correlation coefficient of 0.47, indicating that the mechanisms are dependent of each other. Calcium was found to play a significant role in modulating the effects of stimulus on active hair bundle motility. Blockage of the extrusion pumps that regulate its concentration inside the stereocilia was likewise seen to prolong the induced quiescence, indicating that calcium accumulation may underlie the suppression of spontaneous motility. Research was supported by NIH Grant No. 1R01DC011380.

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**Open and closed-loop gain of the mammalian cochlear amplifier**
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**Background**
The cochlear amplifier is a hypothesized positive feedback process responsible for our exquisite hearing sensitivity. However, experimental evidence for or against the positive feedback hypothesis is still lacking.

**Methods**
In this study, we apply linear control theory to investigate the cochlear amplifier.

**Results**
We determine the open-loop gain and the closed-loop sensitivity of the cochlear amplifier from available measurements of basilar membrane vibration in sensitive mammalian cochleae.

**Conclusion**
We show that the frequency of peak closed-loop sensitivity is independent of the stimulus level and close to the characteristic frequency. This implies that the half-octave shift in mammalian hearing is an epiphenomenon of the cochlear amplifier. The open-loop gain is consistent with positive feedback and suggests that the high-frequency cut-off of the outer hair cell transmembrane potential in vivo may be necessary for cochlear amplification. Acknowledgment: This work is supported by NIH/NIDCD grant R01 DC000141.
In vivo measurement of organ of Corti vibrations near the mouse round-window using phase-sensitive Fourier domain optical coherence tomography

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Background
Hearing loss often has its origin in pathological processes that alter the normal vibratory patterns of the organ of Corti inside the cochlea. Being a mammal and amenable to genetic manipulations, mouse is an excellent animal model to study human hearing and its pathology. However, it has been very challenging to measure the vibration response of the mouse cochlea as the available technology is limited.

Methods
We have developed the phase-sensitive Fourier domain optical coherence tomography (PSFDOCT) and interferometry to image and simultaneously measure the in vivo vibrations of the mouse cochlea.

Results
In this work, we will present the image and in vivo measurements of the mouse organ of Corti vibration at high frequencies (> 50 kHz) viewed through the round-window. Basilar membrane (BM) vibration tuning curve and radial pattern along the BM for high sound levels will be presented.

Conclusion
Simultaneous in vivo measurement of imaging and vibration of organ of Corti in mouse near round window is possible using the PSFDOCT technique. This preliminary study will be improved to study mouse auditory physiology along with genetic modifications. Acknowledgments: This work is supported by NIH/NIDCD grants R01 DC010399 and R01 DC000141.

Evidence Against Power Amplification of the Cochlear Traveling Wave

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Background
The inner ear of mammals mediates sound-induced traveling waves that peak at a frequency-dependent location. It is widely assumed that active processes inject extra power into this wave, thereby enhancing sensitivity, but the power delivered by this cochlear amplification is unknown.

Methods
We determined the power carried by the traveling wave by combining two datasets: (1) We performed two-point interferometric recordings of the gerbil basilar membrane (16-kHz region) in response to tone complexes and derived group delay from the point-to-point phase curves. (2) A recent study by Ren et al. [Nature Commun. 2:216] reported the radial and longitudinal profiles of basilar membrane motion in the 16-kHz region. Using generic wave-physics techniques, we derived from those data the energy density of the traveling wave.

Results
(1) Group velocity increased from 0.9 m/s to 2.1 m/s with sound intensity (10-90 dB SPL). (2) The time average energy per length unit grew from 55 aJ/m to 1.4 pJ/m with sound intensity. (1+2) The power carried by the traveling wave grew from 51 aW to 3.0 pW with sound intensity. For intensities up to 40 dB SPL, the power of the traveling wave at its peak was 1 dB less than the acoustic power entering the middle ear. For higher intensities, the difference grew to 34 dB.

Conclusion
Our findings show that there is no net power gain from middle ear to the peaking traveling wave, and suggest that the nonlinear, compressive response of the cochlea may be realized by variable dissipation rather than saturating amplification.
Cochlear Nonlinearity: The Benefits of Logarithmic Compression

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Background
It has been hypothesized that the medial efferent reflex decreases the cochlear amplifier gain by just the right amount so that the nonlinearity in the basilar membrane response lines up perfectly with the inner hair cell nonlinear transduction process to produce a hair cell receptor potential that is proportional to the logarithm of the sound pressure level (Mountain, AIP Conf. Proc. 1403:238-243.). In this study it is shown that this logarithmic compression provides benefits for coding signals such as speech that have exponential amplitude distributions and also facilitates binaural processing.

Methods
Two theoretical approaches were used. The first approach was to analyze biologically important sounds (e.g. animal vocalizations) for their time-domain statistics and examine the effect of logarithmic compression on coding efficiency. The second approach was to represent signals in the frequency domain using Fourier transforms statistics and examine the effect of logarithmic compression on features related to sound-source localization.

Results
Animal vocalizations (e.g. human speech) often have a probability distribution for amplitude values that is exponential in nature. Logarithmic compression produces a probability distribution that is uniform, a distribution that is optimal for noisey communication channels such as the auditory nerve. The frequency-domain analysis shows that logarithmic compression facilitates the separation of interaural phase and intensity cues. Furthermore it shows that there can be a unique mapping between these interaural cues and the spatial location of a sound source.

Conclusion
The logarithmic compression that is hypothesized to be present at moderate and high sound levels facilitates coding in the auditory nerve and subsequent coding in the medial superior olive complex that contributes to sound-source localization.
Experimental evidence for cochlear amplification and its basis in OHC somatic forces
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Background
Normal mammalian hearing relies on outer-hair-cell (OHC) based electromechanical forces that augment the cochlea’s mechanical response to sound, known as cochlear amplification. Cochlear amplification is frequency- and level-dependent. It amplifies low level sounds more than high level sounds, and the amplification occurs at frequencies close to the local best frequency (BF). Cochlear amplification has been explored theoretically and experimentally. However, the coupling between the OHC electromechanical activity and mechanical responses along the cochlear partition is not well understood.

Methods
We probed cochlear electromechanics with spatially and temporally coincident voltage and pressure measurements using a novel hybrid-sensor positioned close to the sensory tissue. Experiments were performed in anesthetized young adult gerbils. The left pinna was removed and the bulla was widely opened. The basilar membrane (BM) was accessed through a hand-drilled hole in scala tympani (ST) in the basal turn of the cochlea. Compound action potential (CAP) responses to tone pips were recorded at different stages of the experiment to monitor cochlear condition. The hybrid-sensor was constructed in the lab, and was composed of our lab’s standard micro-pressure sensor (OD 125 microns) with a 28 micron diameter, isonel-insulated platinum electrode adhered to its side.

Results
With simultaneous mapping of the intracochlear pressure and extracellular voltage at a series of distances from the BM, BM displacement was derived, resulting in a trio of coincident pressure, voltage, and displacement measurements. We found a phase shift between voltage and displacement, approximately half an octave beneath the peak frequency, which would allow power injection from OHC somatic forces. Moreover, at low SPL the phase between pressure and displacement indicated negative resistance, signifying that biomechanical elements within the sensory tissue were amplifying the power in the traveling wave.

Conclusion
These results are the most concrete evidence for intracochlear power amplification to-date and support OHC somatic forces as its source.
Power Flow Calculation in the Cochlea by Estimation of the Spatial Patterns of Intracochlear Pressure and Basilar Membrane Response

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Background
Power flow in the cochlea is an important indicator of the presence (or absence) of energy generation from active process, the so-called cochlear amplifier. Calculation of spatial power flow requires the knowledge of both the basilar membrane (BM) velocity and the intracochlear pressure in the spatial domain. However, due to technical difficulties, direct measurements of these two quantities with fine spatial resolution are not available. The purpose of this study is to calculate the power flow in the cochlea based on available experimental data.

Methods
The spatial pattern of the BM velocity is estimated from the BM spectrum response by using the frequency-location mapping. Measured BM spatial response around a best-frequency (BF) place is used to adjust the transformation by using the traditional frequency-location mapping. The intracochlear pressure is estimated from both the WKB method and the finite element method (FEM) as a postprocessing step, using the measured (near the BF place) or estimated (over a larger spatial range) BM velocity as the input. The accuracy of the pressure estimation is checked by comparing the application of the algorithm to the self-consistent FEM data, using FEM predicted BM velocity as the input. The power flow in the cochlea is calculated at each cross section along the longitudinal direction.

Results
The developed algorithm predicts the wavenumber as a function of the longitudinal coordinate near the BF place. The accuracy of predicted pressure depends on the spatial extent of the known BM velocity. Knowledge of the BM velocity basal to the BF place is found to be required for accurate power estimation.

Conclusion
The power flow in the cochlea decays longitudinally if the cochlea is passive. In an active cochlea, the power flow increases significantly around the BF place.
Intracochlear pressures with bone conduction stimulation in human temporal bones
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Background
Experimental studies on bone conduction on temporal bones or anatomical whole heads have often measured the velocity or acceleration of the promontory as an estimate for the cochlear input. The aim of this study is to measure pressures in cochlear scalae to investigate the mechanism of bone conduction on temporal bones. Measurements of intracochlear pressures during sound stimulation and round-window stimulation have provided important information. Similarly, the differential pressure across the partition – which estimates the cochlear drive at the base of the cochlea – can be measured during bone conduction, providing information not previously available.

Methods
Custom-made fiberoptic pressure sensors (diameter of 200 $\mu$m) were used to simultaneously measure the intracochlear pressures of the scala vestibuli and the scala tympani in human temporal bones. After insertion of the sensors (100 $\mu$m deep) through cochleostomies, the sensors were sealed to the surrounding bone with dental impression material (Jeltrate) followed by dental cement to firmly fix the sensors to the cochlear promontory, minimizing relative movement of the sensors with respect to the bone. Holders for the fiberoptic cables were released to remove significant friction along the cable to allow them to vibrate freely. The bone was stimulated with pure tones between 100-20000 Hz using a commercially available bone anchored hearing system at a constant electrical voltage (1 V peak).

Results
During bone conduction stimulation, the relative movement of the fiberoptic sensors near the cochleostomies versus bone was generally below 5 dB, thus the sensors were vibrating similarly to the cochlea. Intracochlear pressures with signal to noise ratios of more than 10 dB and linear were recorded for frequencies between 300 and 8000 Hz. In the same frequency band, the magnitude of the differential pressure during bone conduction stimulation was in the same order of magnitude as the differential pressure obtained with ear-canal sound stimulation of 1 Pa.

Conclusion
With modifications of our method for intracochlear pressure measurements, we can now understand the effect of bone conduction on scalae pressures and the cochlear drive in human temporal bones.
INTRACOCHLEAR PRESSURE MEASUREMENT USING COMMERCIALLY AVAILABLE SENSORS

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Background
Measurement of differential intracochlear pressure has proven to be a useful technique for the exploration of normal and pathological processes of ossicular transmission and the effectiveness of stimulation through bone conduction and the round window. Previous studies have employed custom-fabricated micro-optical pressure sensors with special purpose signal-conditioning circuitry, or have taken measurements at two different sound pressure levels so as to overcome the dynamic range limitations of commercially available sensors.

Methods
In this study, a system using off-the-shelf fiber optic pressure sensors and signal conditioners was used for measurement of intracochlear pressures in human cadaveric temporal bones concurrently with laser Doppler vibrometry (LDV) measurement of ossicular and round window velocities at the same sound pressure levels. Measurements of pressures in the scala vestibuli (SV), scala tympani (ST) and external auditory canal (EAC) were made concurrently with LDV measurement of either stapes or round window velocity.

Results
Experimentally obtained estimates of SV/EAC and ST/EAC transfer functions, middle ear gain and group delay, cochlear fluid volume velocity and cochlear impedance were taken for a sample of normal human temporal bones and found to be consistent with previously published results.

Conclusion
The technique of using off-the-shelf fiber optic pressure sensors and signal conditioners offers a significant reduction in the cost and difficulty of intracochlear pressure measurement and may prove useful in the further exploration of normal and pathological auditory mechanics, assessment of performance in acoustic implants, and modeling of acoustic trauma.

Local Transfer Characteristics of the Basilar Membrane

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Background
Studies on cochlear mechanics mainly report motion of single basilar membrane (BM) locations. However, comparison of responses from two locations in the same preparation can give much additional information on the propagation of the traveling wave. In a recent study, Ren et al. [2011, PlosOne 6:e20149] reported two-point BM recordings and introduced "local transfer functions," obtained by dividing the complex spectra of the responses at the two points to identical stimuli. This cancels out any irregularities in calibration and/or middle-ear transfer. In this study, we used local transfer functions to analyze the propagation of the traveling wave.

Methods
A single-point laser vibrometer measured the motion of two beads placed (typically <400 µm apart) on the BM in anesthetized Mongolian gerbil. We stimulated with irregularly-spaced, equal-amplitude tone complexes of varying sound intensity. The responses of the two beads to identical stimuli were used to construct local transfer functions.

Results
(1) Phase velocity and wavelength of the traveling wave were strongly intensity dependent, confirming the findings of Ren et al. (2) Phase velocity was minimal for frequencies just below the characteristic frequency (CF) and did not decrease further with increasing frequency. (3) Above CF, bead-to-bead amplitude and bead-to-bead phase both steadily decreased with frequency; their relative rate was approximately 54 dB/cycle.

Conclusion
(1) The strong dependence of phase velocity on intensity suggests that damping is not the only mechanical property of the cochlear partition that changes with intensity. Most likely, the effective stiffness of the BM changes with intensity, too. (2)
The absence of a continued decrease of phase velocity with frequency, even for frequencies well above CF, indicates that resonance does not play a role in the frequency selectivity of the BM. (3) The ratio of amplitude and phase decrease observed above CF contradicts the mass dominated motion expected from resonance, and is consistent with a damping dominated motion expected from the assumption that the cochlear partition has negligible fixed mass.

Transmission of distortion products in the mammalian cochlea

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Background

Otoacoustic emissions (OAEs) are sounds originating in active cochlear mechanics that travel back out to the ear canal (EC). The mechanism by which OAEs are transported out through the cochlea is not certain. In most cochlear models, the cochlea was assumed to operate similarly in forward and reverse propagation and the OAE was supposed to travel out of the cochlea via a reverse traveling wave along the cochlear partition. Another possibility is that reverse transmission is mediated via a compression pressure in the cochlear fluid. Past experimental observations have implicated both these reverse transmission modes. To probe the role of cochlear fluid in reverse transmission, two-tone stimulation was used and the distortion products (DPs) generated in the cochlea were considered to be the intracochlear sound sources. The DPs drove the stapes in reverse and were detected in the EC as distortion product otoacoustic emissions (DPOAEs). Reverse transmission was characterized by comparing intracochlear pressure responses at the basilar membrane (BM) and at the stapes.

Methods

Experiments were performed in anesthetized young adult gerbils. The left pinna was removed and the bulla was widely opened. The BM was accessed through a hand-drilled hole in scala tympani (ST) ~ 2.2 mm from the base (~20% of the cochlear length). Simultaneously, the intracochlear pressure responses were measured in scala vestibuli (SV) at the very base, next to the stapes. Compound action potential (CAP) responses to tone pips were recorded at different stages of the experiment to monitor the cochlear condition.

Results

Comparison between the ST and SV pressures at primary and DP frequencies allowed us to characterize sound transmission in the forward and reverse directions. By comparing the phases of the ST and SV pressure the travel delay between the two locations was determined, and reverse transmission was found to be consistent with a reverse traveling wave. In addition, transverse spatial variations in ST pressure were similar at primary and DP frequencies. Because the spatial variation reflects traveling wave wavelength, this similarity reinforced the notion that reverse propagation was mainly via a traveling wave mode along the cochlear partition, rather than via a compression pressure through the cochlear fluid.

Conclusion

Sound reverse propagation is primarily via a reverse traveling wave, rather than the cochlear fluid.
Making Waves in the Tunnel of Corti
Aleks Zosuls¹, Laura Rupprecht¹, David Mountain¹
¹Boston University

Background
Evidence of fluid flow in the tunnel of Corti (ToC) was demonstrated by Karavitaki (Biophysical Journal, vol.92, p.3284-3293). Experiments by Chen et. Al. (Nature Neuroscience, vol.14, p.770-774) show that the basilar membrane (BM) and reticular lamina don't always move in phase. This differential motion could create a local pressure in the fluid filled ToC. These results indicate that fluid flow in the ToC may be significant for tuning and coupling in the hearing organ.

Methods
Experiments were performed on fresh excised gerbil cochleae. The scala vestibuli and tympani were opened to access and image the organ of Corti (OC). A blunted micropipette pushed on the BM under the outer pillar cell to create a point pressure. The pipette was driven by a piezoelectric stack. Radial motion of the inner hair cell stereocilia hair bundles (IHCSS) and lateral motion of the tunnel crossing fibers (TCF) in the ToC was observed by stroboscopic imaging under high magnification. The motion of IHCSS and TCF bundles was quantified using a cross correlation algorithm.

Results
When the probe was indenting the BM, the TCF fibers moved longitudinally (basal or apical) away from the probe. Assuming the TCF’s move with the flow of fluid in the ToC, the TCF motion can be interpreted as flow in the tunnel. There was a 180 degree transition at the probe location signifying the fibers moving in opposite directions (figure1). When moving away from the probe basal or apical there was a building phase delay in the fiber motion that is evidence of traveling wave propagation (figure1). The propagation velocity was estimated via curve fitting to be 2.3 m/s. The fit ignored data points near the probe to eliminate the transition region.
When the probe was indenting the BM the IHCSS moved toward the modiolus due to motion of the OC. Similar to the TCF phase, the phase of the IHCSS had a building delay moving away from the probe (figure1). This is evidence of a travelling wave with a propagation velocity estimated to be 2.2 m/s.

Conclusion
There is no evidence that the pillar cells move longitudinally during the probe stimulation which indicates that the TCF’s are driven by fluid flow, not movement of their anchor points. The similar traveling wave velocities in both structures indicate that there is coupling between IHCSS motion and fluid flow in the ToC.
Assessment of superior canal dehiscence location and size on intracochlear sound pressures

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Background
Superior canal dehiscence (SCD) is a defect in the bony covering of the superior semicircular canal. Patients with SCD syndrome present with a wide range of hearing loss. Some clinical studies have suggested that an increase in SCD size is associated with increased hearing loss, while others have observed no association. Our previous human cadaveric temporal bone study showed that increase in dehiscence size (up to 2 mm long and 0.7 mm wide) causes a low-frequency monotonic decrease in the cochlear drive across the partition, consistent with increased hearing loss. At stimulus frequencies of more than 1 kHz, a small dehiscence (0.5 mm diameter) sometimes produced a larger decrease in cochlear drive than larger dehiscences. But the effects of larger SCDs (> 2mm long) were not studied, although most SCD patients have larger than 2 mm-long SCDs. Furthermore, it is unknown whether hearing is affected by the location of the SCD. In this study we assess the effect of SCD location and the effect of large sized SCD on intracochlear pressures.

Methods
We used simultaneous measurements of intracochlear sound pressures in scalavestibuli and scala tympani to determine the sound-pressure difference across the cochlear partition, which is a measure of hearing loss in a temporal bone preparation. We measured the pressure difference before and after SCDs at different locations (e.g. closer to the ampulla of the superior semicircular canal or closer to the common crus), and for different dehiscent sizes (including larger than 2 mm long and 0.7 mm wide).

Results
Our measurements suggest: 1) Changes in SCD location do not affect the intracochlear pressure difference. 2) Larger SCDs produce larger decreases in the pressure difference at low frequencies. However, the effect of SCD size seemed to saturate as the size increased above 2-3 mm long and 0.7 mm wide. Although these trends were generally consistent across ears, the amount of pressure change due to dehiscence varied across ears.

Conclusion
These findings show that SCD size can help explain some of the variability of hearing loss in patients and that the location of the SCD does not influence the amount of hearing loss. However, different ears varied in the extent of hearing loss for similar SCD sizes.
Decreasing Stiffness Reduces Spread of Excitation via Tectorial Membrane Waves

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Background
Classical models have treated the tectorial membrane (TM) as a resonant system that can increase sensitivity and sharpen cochlear tuning (Allen, 1980). Recently, it has been shown that the TM supports longitudinal traveling waves (Ghaffari et al, 2007) and that the spatial extent of these waves is strongly correlated with cochlear tuning (Ghaffari et al, 2010). Here we investigate the relationship between stiffness and tuning for TM waves and compare this relation to predictions from resonance models.

Methods
We measured wave and material properties of isolated mouse TMs equilibrated at pH 7 and 4. Wave decay constants and speeds were measured by stimulating suspended TM samples at acoustic frequencies (Ghaffari et al, 2007). TM stiffness and viscosity were measured using microfabricated shear impedance probes (Gu et al, 2008).

Results
Reducing bath pH from 7 to 4 decreased shear stiffness (45% change) but had a minor effect on shear damping, as predicted by gel models of the TM (Freeman et al, 1997). Wave measurements at normal and acidic pH showed that the spatial extent of TM waves decreased by 42% but wave speed changed relatively little (<12%). These findings suggest that decreasing shear stiffness (without changing viscosity) reduces spread of excitation via TM waves.

Conclusion
Our results show that decreasing stiffness reduces spread of excitation in TM waves. By scaling symmetry, this change would tend to sharpen cochlear tuning. In contrast to previous models, this sharpening occurs without changes in viscous damping in the subtectorial space (Allen, 1980) or changes in outer hair cell motility (Neely, 1993). These findings show that TM wave and resonance theories differ significantly. While both suggest that the TM plays an important role in determining cochlear tuning, the underlying mechanisms are completely different.
In vivo vibrometry at the apex of the unopened mouse cochlea
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Background
The study of cochlear mechanics has provided important insights into cochlear function. Previous experiments from the base of the opened, mammalian cochlea have shown frequency selectivity and non-linear compression. However, in vivo measurements of cochlear mechanics have traditionally been difficult to obtain at the apex of the mammalian cochlea. Recent advancements in imaging technology provide the opportunity to make such measurements. Herein, we present in vivo images and measurements from the apex of unopened mouse cochlea using spectral domain optical coherence tomography (OCT).

Methods
Spectral domain OCT is a non-invasive interferometric imaging technique that generates images based on depth-resolved sample reflectivity through the use of a fast Fourier transform. By analyzing the phase of the reflected signal at each depth, OCT can also be used as a vibrometer. Using OCT, we were able to visualize and measure organ of Corti vibrations approximately a half turn from the mouse cochlear apex. At the lowest stimulus intensity, we averaged to ensure a noise threshold <1nm, defined as the mean of the noise plus three times the standard deviation of the noise.

Results
The characteristic frequency (CF) of the locations we recorded from was between 7 to 10 kHz. We found that compression was highest at the CF with growth rates of ~0.2 dB/dB. The Q10-dB value at a stimulus level of 50 dB SPL was ~2. These values are similar to previously measured values from the cochlear base in other species with CFs of between 6 to 18 kHz.

Conclusion
Our results suggest that the cochlear amplifier functions in the mouse apex much like it does in the base of larger mammals.

Handling of experimental data on the cochlea obtained with Optical Coherence Tomography (OCT).
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Background
With the technique of Optical Coherence Tomography (OCT) it is possible to measure movements of the basilar membrane (BM) and the Reticular Lamina (RL) separately. In the dead animal these movements were found to be nearly identical. In the living animal, in a viable cochlea, the RL moves in the region of maximum response with an amplitude about three times larger and a phase difference less than 90 degrees.

Methods
This implies that for the intact cochlea we have to take into account a definite oscillatory movement of the fluid inside the Organ of Corti (OoC) and the Internal Spiral Sulcus (ISS). In a most simplified and perhaps exaggerated way it can be stated that ‘activity’ of Outer Hair Cells (OHCs) operates on this mass of fluid.

Results
It will be shown how this problem can be treated in cochlear modeling. Specific experimental evidence on lateral fluid motion lacking, we have to follow paths defined by assumptions about the movements of the fluid contained in those two spaces.

Conclusion
The consequences of various sets of assumptions will be demonstrated.
With the technique of Optical Coherence Tomography (OCT) it is possible to measure movements of the basilar membrane (BM) and the Reticular Lamina (RL) separately. In the dead animal these movements were found to be nearly identical. In the living animal, in a viable cochlea, the RL moves in the region of maximum response with an amplitude about three times larger and a phase difference less than 90 degrees. This implies that for the intact cochlea we have to take into account a definite oscillatory movement of the fluid inside the Organ of Corti (OoC) and the Internal Spiral Sulcus (ISS). In a most simplified and perhaps exaggerated way it can be stated that ‘activity’ of Outer Hair Cells (OHCs) operates on this mass of fluid. It will be shown how this problem can be treated in cochlear modeling. Specific experimental evidence on lateral fluid motion lacking, we have to follow paths defined by assumptions about the movements of the fluid contained in those two spaces. The consequences of various sets of assumptions will be demonstrated.

Lgr5-positive cells contribute to hair cell regeneration in the damaged mouse utricle

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Background
Hearing loss and balance disturbance are sensory disorders affecting millions of patients, with sensory hair cell (HC) loss being a primary pathology in both disorders. Previous studies illustrated that the postnatal mouse utricles contain cells capable of regenerating HCs after pathologic HC loss, yet our understanding of their exact identity and regulatory mechanisms is limited. Wnt signaling regulates cellular repair in several organ systems. Here, we hypothesize that the Wnt target gene Lgr5 marks HC precursors in the damaged mouse utricle.

Methods
Utricles from embryonic (E) and postnatal (P) mice were examined for Lgr5 expression. To induce HC loss, P3 mice were cultured with neomycin (1.0mMx24hr) and then in drug-free media for 2 to 13 days. Lgr5EGFP-CreERT2/+ transgenic mice or quantitative PCR were used to determine Lgr5 expression. To fate-map Lgr5+ cells, a cre-loxP approach using the Lgr5EGFP-CreERT2/+; Rosa26RtdTomato/+ mice was taken. Time-lapse imaging of fate-mapped cells in cultured utricles using spinning disk confocal microscopy was performed. After culture, tissues were fixed and immunostained for markers of HCs (Myosin7a), supporting cells (Sox2 and Plp1), and cell death (TUNEL).

Results
In the Lgr5EGFP-CreERT2/+ mice, EGFP expression was found in HCs and supporting cells at E15.5 and 18.5, then became undetectable by P3. After neomycin treatment, Lgr5-EGFP was upregulated in supporting cells in the striolar region, where HC loss concentrated, and subsequently persisted for at least 6 days. Quantitative RT-PCR also found Lgr5 upregulation in neomycin-treated wildtype utricles. Two days after damage, myosin7a+ HCs decreased by 56% and Lgr5+ cells (70.3±9.2 per 20,000 µm²) were 99.8% Myosin7a-, 99.4% Sox2-, 100% Plp1+, and 97% TUNEL-. Five days after damage, lineage tracing of Lgr5+ cells showed that 84.7% of traced, tdTomato+ cells (48.0 per organ) were myosin7a-residing in the supporting cell layer. tdTomato+, myosin7a+ cells significantly increased from 5 to 12 days after damage (7.3±2.5 to 17.7±7.0 per organ). Time-lapse imaging from 2.5-6 days after damage demonstrated that tdTomato+ cells underwent dynamic morphological changes, a subset of which was reminiscent of nascent HCs.

Conclusion
We found Lgr5 expressed in the embryonic utricles and downregulated at birth. HC loss reactivates Lgr5-EGFP expression among supporting cells that were fate-mapped to regenerated HC-like cells. Together our data suggest that Lgr5+ cells behave as facultative HC precursors.
Regeneration of Mammalian Cochlear and Vestibular Hair Cells through Hes1/Hes5 Modulation with siRNA
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Background
The Notch pathway is a cell signaling pathway determining initial specification and subsequent cell fate in the inner ear. Previous studies have suggested that new hair cells (HCs) can be regenerated in the inner ear by manipulating the Notch pathway.

Methods
The organ of Corti and vestibular maculae of postnatal day 3 (P3) mouse pups were harvested and cultured in vitro. siRNA to Hes1 and Hes5 using a transfection reagent or siRNA to Hes1 encapsulated within poly(lactide-co-glycolide acid) (PLGA) nanoparticles were added to cultures.

Results
Increased HC numbers were observed in the cultured and untreated cochleae and maculae, and in the cultured cochleae and maculae pre-conditioned with a HC toxin (4-hydroxy-2-nonenal or neomycin) and then treated with the various siRNA formulations. A decrease of Jagged1 positive supporting cells (SCs) was observed in cultured cochleae of mouse pups treated with Hes1 siRNA and transfection reagent. Treating cochleae with siRNA to Hes1 associated with a transfection reagent or siRNA to Hes1 delivered by PLGA nanoparticles decreased Hes1 mRNA and up-regulated Atoh1 expression allowing SCs to acquire a HC fate. Experiments using cochleae and maculae of p27kip1-GFP transgenic mouse pups demonstrated that newly generated HCs trans-differentiated from SCs. PLGA nanoparticles are non-toxic to inner ear tissue, readily taken up by the tissues, and present a synthetic delivery system that is a safe alternative to viral vectors.

Conclusion
These results indicate that when delivered using a suitable vehicle, Hes siRNAs are potential therapeutic molecules that have the capacity to regenerate new HCs in the inner ear and possible restore human hearing and balance function.

Preparing the Cochlea for Stem Cell Implantation
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Background
Technology for growing hair cells from stem cells (SCs) in vitro is rapidly advancing. In parallel, it is necessary to design methods for inserting SCs into the auditory epithelium (AE) of deaf ears devoid of hair cells (HCs), in vivo. Inserting cells into an epithelial cell layer is complicated in any organ, and is even more complex in the hostile, high potassium environment of the scala media (SM). The robust junctional complexes between cells in the AE present an additional obstacle to SC integration. A successful therapy will require methods that optimize survival of exogenous cells injected into SM and their integration into the AE by making the cochlear environment more tolerable and relaxing intercellular junctions. To that end, we evaluated the survival of cells in artificial cochlear fluids in vitro and then manipulated the cochlear environment in vivo to open intercellular junctions and increase its hospitability to exogenous cells.

Methods
Cell survival in cochlear fluids was evaluated by monitoring fluorescence of mCherry labeled HeLa cells after adding artificial fluids (endolymph, perilymph or endolymph+perilymph) to the culture. In vivo experiments used neomycin-deafened guinea pigs. To render SM more hospitable to exogenous cells, we reduced potassium levels by (a) injecting furosemide, and (b) replaced the endolymph with artificial perilymph. To disrupt the adherens junctions in the AE we injected sodium caprate via a cochleostomy. We then injected HeLa cells into the SM. Changes of the AE junctions were evaluated in whole-mounts of the AE stained for ZO-1. Presence of HeLa cells in the cochlea was evaluated by fluorescence stereomicroscopy followed by epi-fluorescence on whole-mounts of AE.

Results
Cells exposed to artificial endolymph detached from the dish and became spherical promptly. Most cells died within six hours. Cells cultured in perilymph or in the endolymph-perilymph mixture survived. In the AE of sodium caprate-treated
ears, gaps appeared between cells, suggesting deterioration of apical junctions. One day after the injection, implanted exogenous HeLa cells were observed in the AE and other areas in ears treated with sodium caprate and furosemide.

**Conclusion**
Transient manipulations of the cochlear environment facilitate survival of exogenous cells for at least one day. Further refinements are needed to prepare the deaf cochlea for SC therapy.

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**Age-related alterations in neural crest-derived cells of the mouse inner ear**

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¹Medical University of South Carolina

**Background**
Glial cells in the auditory nerve and cells in the cochlear lateral wall undergo turnover in response to several stress conditions, although their ability to repopulate themselves declines with age. The neural crest is a transient embryonic structure that gives rise to a wide variety of cell types including glial cells, some neurons of the peripheral nervous system and mesenchymal cells. Multi-potent neural crest-derived stem cells (NCSCs) have been found in several adult tissues and organs suggesting an important role of the neural crest-derived cells in the maintenance of adult tissue homeostasis and self-repair after injury. This study seeks to correlate the effects of aging on the self-renewal of neural crest-derived cells in the adult inner ear with age-related alterations in gene expression profile.

**Methods**
The sphere formation assay was used to determine the relative numbers of NCSCs in the auditory nerves and cochlear lateral wall of young adult (1-3 mo) vs. aged (18-30 mo) CBA/CaJ mice. Gene expression profiles of cochlea from young adult vs. middle to early old aged (12-16 mo) mice were examined using gene microarray, proteomic analysis, western blot and quantitative immunohistochemistry. Twelve to 16 month old mice were used to determine changes in gene expression that might precede significant cell loss that occurs in old age.

**Results**
Our data show that cells isolated from the auditory nerves and cochlear lateral wall tissues of both young and old adult mice formed spheres in culture that displayed extensive self-renewal capacity and contained cells that express p75 and Sox10, markers of a neural crest lineage. Application of ouabain, an inhibitor of Na+/K+-ATPase, into the cochlea significantly increased the number of NSCSs from the auditory nerves of young adult mice. This was not the case with auditory nerves of older mice. Gene expression profiles of cochlea from young adult vs. middle to early old aged mice revealed age-related changes in expression of more than 40 genes associated with the regulation/differentiation of neural crest-derived cells.

**Conclusion**
Age-dependent hearing loss and pathological changes in the inner ear may be due to a decline in the number of neural crest-derived cells suggesting an age-related loss of self-renewal capacity. Expression profiling studies revealed potential targets for restoring cell renewal in the adult inner ear. This study was supported by NIDCD R01DC7506, P50DC00422 and NCRR P20RR16434.
Molecular Characteristics of Severely Damaged Organ of Corti
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Background
Humans with severe or profound hearing loss have cochlear regions where putative support cells remain despite total hair cell loss, as well as regions where the organ is replaced with a flat unspecialized epithelium (Teufert et al., 2006; Hoa et al., 2010 ARO). Understanding the characteristics of the cells that remain after hair cell loss will be crucial to identifying feasible regenerative procedures.

Methods
Adult C57BL/6J mice were damaged by kanamycin/furosemide injection (Oesterle et. al., 2008), and transgenic mice that express diphtheria toxin (DT) receptor under control of the HC-specific promoter pou4f3 were injected with DT to kill hair cells. Mice were killed 8 days or 2 months later, and whole-mount preparations of the organ of Corti were immunohistochemically processed using support cell (acetylated tubulin, Kir 4.1, monocarboxylate transporter 2 (MCT2), Sox2) and sulcus cell markers (CD44).

Results
With both damage paradigms, regions are seen where putative support cells remain despite total hair cell loss. Sox2, MCT2, and acetylated tubulin immunolabeled cells in these regions, and CD44 was expressed in cells in the flanking inner and outer sulcus. Regions of flat epithelium were seen in the C57BL/6J mice 2 months after kanamycin/furosemide administration. In the apical and middle turns of these animals, patches of flat epithelium were interspersed between segments of damaged epithelium containing pillar and Deiters’ cells (acetylated tubulin+/Sox2+ cells). Fewer nuclei are present in the flat epithelium and these cells express CD44, but not Sox2 or acetylated tubulin.

Conclusion
1) Differentiated support cells can survive despite total hair cell loss and retain at least some cellular identity, and 2) Sulcus cells may migrate into the region formerly occupied by the organ of Corti.

Supported by the Hearing Heath Foundation and NIDCD grants R01 DC03944 and P30 DC04661.
Deciphering the strategy of hair cell regeneration in zebrafish lateral lines
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Background
The capacity and capability of hearing restoration upon hair cell impairments in the ear are apparently species dependent, even though the hair cells are killed essentially in the same way. For instance, cisplatin, an anti-cancer drug, frequently induces hearing loss in patients and kills zebrafish hair cells of inner ear and lateral lines; however, only fish can regenerate the hair cells after the treatment is terminated. Therefore, we ask why and how regeneration program is triggered in the neuromasts, which cells within the neuromasts become the nascent hair cells, and how these cells decide to undergo proliferation vs. differentiation.

Methods
Transgenesis in zebrafish; in situ hybridization; immunohistochemistry; small chemical screen; live imaging.

Results
We have observed that neuromast hair cells undergo apoptosis during the cisplatin treatment, and the damaged neuromasts start a slow and steady regeneration meanwhile. Interestingly, when the cell death, induced by other reagent(s), is not apoptotic, the hair cell regeneration process is quicker; suggesting different types of hair cell death may result in different ways to regenerate. Molecularly, the cisplatin treatment enables higher expression of Atoh1 and Sox21 in the sensory and periphery support cells, respectively, and a rapid proliferation in a group of periphery non-hair cells. Furthermore, through small chemical compound screens of hair cell regeneration regulators, we've obtained 4 inhibitors and determined in which step(s) these compounds act.

Conclusion
Our study has laid a ground to explore the mechanism(s) underlying zebrafish lateral line hair cell regeneration.
The role of planar cell polarity pathway in the ectopic regenerated hair cells regulated by the dynamic ADF/destrin expression

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Background
Planar cell polarity (PCP) signaling regulates cochlear extension and coordinated orientation of sensory hair cells in the inner ear. Studies have shown retrovirally-mediated introduction of Atoh1 transcription factor is capable of causing some mature supporting cells to transdifferentiate into hair cells. ADF/Destrin is one important factor in actin depolymerization families, which is essential for several cellular processes including cell survival, shaping, cytokinesis and migration. However, the role of PCP pathway in ectopic regenerated hair cells regulated by dynamic ADF/destrin expression is still unknown.

Methods
Ad5-EGFP-math1 was used to over-express math1 in the cochlea of neonatal mice in vitro. We compared dynamic distribution of ADF/destrin between developing cochlear and the ectopic regenerated hair cells, investigated the establishment of ectopic hair cell polarity and the relationship with ADF/destrin expression. We also observed the proliferation, differentiation and epithelial PCP in the lesser epithelial ridge (LER) down-regulated ADF/destrin after testosterone-BSA treatment in vitro.

Results
It showed that ADF/destrin distributed in some cuticular plate’s border of both hair cells and supporting cells at E14.5, then expressed in the whole cuticular plate of hair cells and supporting cells at E18.5 and P1, there were no ADF/destrin in the hair cells after P4. Some ectopic regenerated Myo7a(+) cells expressed ADF/destrin within one week and disappeared 10 days after over-expressing Atoh1, which is compared with the high level differentiated ectopic hair cells. After ectopic regenerated Myo7a(+) cells developed actin-rich stereocilia, their basal body moved from center to the distal side, suggesting the underlying PCP establishment in ectopic regenerated myo(+) cells. The level of ADF/destrin decreased after testosterone-BSA treatment 12-24h in vitro, more BrdU(+) cells and Myo7a(+) cells were observed in LER, however, the CE and cell polarity of Ad5-EGFP-math1 infected LER is not affected obviously.

Conclusion
Our results indicate that ADF/destrin participate in the development of sensory epithelia in mammalian cochlear and ectopic hair cells induced by over-expressing Atoh1. PCP signaling exists in the development of ectopic hair cells. Together, these data suggest that ADF/destrin dependent actin depolymerization maybe not essential for hair cell polarity while it is required for cell proliferation and differentiation of cochlear LER regions.

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Characterization of Human Pluripotent Stem Cell Derived Otic Progenitor-like Cells

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Background
Hearing loss is primarily caused by loss of sensory hair cells (HCs) in the organ of Corti within the cochlea. Although human pluripotent stem cells are a promising source for cell therapy, there are few reports of the efficient generation of otic progenitor or HC-like cells from human pluripotent stem cells. The auditory system develops alongside the central nervous system and is dependent for proper formation upon the signaling from the developing adjacent neural ectoderm. This study tested the hypothesis that when directing human pluripotent stem cells towards a neural fate, a sub-population of the cells will become otic progenitor cells. This work is a continuation of prior investigations using a well-characterized neural differentiation paradigm to direct human embryonic and induced pluripotent stem cells towards an otic progenitor cell fate.

Methods
Human H9 ES cells and IMR90-4 iPS cells were differentiated using a well established neural differentiation paradigm. An otic progenitor-like population of cells was characterized using immunocytochemistry, RT-PCR and qPCR analysis.

Results
Neural differentiation of human pluripotent stem cells identified a distinct subpopulation of cells expressing markers of an otic progenitor fate. The gene transcript and the protein expression profile were consistent with that seen during early mammalian inner ear development. Furthermore, the sequence and expression pattern of marker transcripts during differentiation of human pluripotent stem cells towards an inner ear lineage in vitro appears similar to that seen during normal mammalian inner ear development. In addition, we have found that modification of the neural differentiation protocol guided by knowledge of early mammalian inner ear developmental signaling mechanisms reliably leads to an enrichment of this otic progenitor-like population. Ongoing studies aim to further optimize the culture paradigm to generate a more robust otic progenitor cell population for use in development and disease modeling as well as transplantation-based studies.

Conclusion
Neural differentiation of both embryonic stem cells and induced pluripotent stem cells leads to the expression of a panel of marker transcripts characteristic of developing otic progenitor cells as analyzed by immunocytochemistry, RT-PCR and qPCR analysis.
Characterization of canonical Wnt/β-catenin signaling activity in the developing and adult mouse cochlea

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Background
During development canonical Wnt/β-catenin signaling plays important roles in organogenesis by regulating proliferation and differentiation. In more mature tissues this pathway functions in progenitor and stem cell maintenance. Lgr5, a downstream target of Wnt/β-catenin signaling, is expressed in the embryonic and mature organ of Corti, and recent studies have identified a role for canonical Wnt signaling during cochlear sensory epithelium development. However, the pattern of direct Wnt/β-catenin activity had yet to be characterized in adult cochleae, thus we utilized a transgenic reporter mouse which provides highly specific single-cell-resolution readouts of canonical Wnt activity to compare signaling in developing and adult cochleae.

Methods
We analyzed Wnt/β-catenin activity patterns using cochleae from TCF/Lef:H2B-GFP reporter mice which possess six copies of a TCF/Lef responsive element linked to an hsp68 promoter that drives expression of a nuclear H2B-GFP fusion protein. Embryonic cochleae from stages E12-E17.5 and adult cochleae from 8-10-week-old mice were collected, and analyzed by immunohistochemistry (n>4 cochleae analyzed per stage). Tissue was co-immunostained with antibodies against the support cell/progenitor cell marker Sox2, and either the proliferation indicator Ki-67 or the hair cell marker MyosinVI.

Results
During the early proliferative phase of development, nuclear GFP activity was observed throughout the floor of the cochlear duct where in some cells it was co-expressed with the prosensory marker Sox2 and/or the mitotic marker Ki-67. At E14.5, during the onset of hair cell differentiation, the level of GFP expression was significantly reduced. By E17.5, when HC differentiation is mostly complete, low levels of GFP were detected within support cells of the organ of Corti, and within the GER. In adult cochleae, GFP expression could still be detected within a subset of cells within the organ of Corti.

Conclusion
In cochleae from TCF/Lef:H2B-GFP reporter mice, Wnt/β-catenin activity appeared highest in younger prosensory cells and diminished with the onset of differentiation, with some activity being maintained through adulthood. The early pattern of activity was similar to that previously described in the Lgr5 reporter mice, demonstrating the cochlear specificity of the TCF/Lef:H2B-GFP reporter. A solid understanding of the progressive changes in Wnt/β-catenin signaling may provide valuable insight for identifying cells which may serve as targets for inducing regeneration in damaged adult cochleae.
Bone Morphogenetic Protein and Inhibitor of Differentiation Antagonize Hair Cell Differentiation during Avian Hair Cell Regeneration in Post-Hatch Chickens

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Background
The transcription factor Atoh1 is both necessary and sufficient to induce hair cell production during development. During avian hair cell regeneration, however, neither endogenous Atoh1 activity nor overexpression of Atoh1 fully predict hair cell fate. In developing auditory epithelia, bone morphogenetic proteins (BMPs) antagonize the expression of Atoh1 via another set of transcription factors, Inhibitors of Differentiation (IDs).

Methods
To begin to explore the role of BMPs and IDs in auditory hair cell regeneration, we performed in situ hybridization to examine expression of these genes in normal and Streptomycin-damaged basilar papillae (BPs). We also tested the effect of inhibiting or activating BMP signaling in damaged BPs on 1) the levels of Atoh1 and Id transcripts and 2) the numbers of regenerated hair cells. Additionally, in order to test the hypothesis that IDs antagonize hair cell differentiation, we electroporated full-length Id3 or control plasmid into supporting cells in cultured BPs and examined effects on the numbers of supporting cells that transdifferentiated as hair cells.

Results
In situ hybridization showed that control basilar papillae have high expression of Id2, Id3, and Bmp4 transcripts and very low levels of Atoh1 and Bmpr1a (Bmp2/4 receptor) transcripts. By 1 day post-Streptomycin, Atoh1 and Bmpr1a are upregulated in the damaged area, whereas Id2, Id3, and Bmp4 are downregulated. Treating tissue with BMP2 results in decreased Atoh1 transcript, and decreased Id2 and Id3 transcripts. Conversely, treating with noggin, a BMP inhibitor, results in decreased Id2 and Id3 transcripts. Accordingly, the number of hair cells that differentiate increases with noggin treatment, while the number of hair cells that differentiate decreases with BMP4 treatment. Preliminary analysis indicates that overexpression of Id3 in supporting cells results in significant reduction of supporting cell transdifferentiation into hair cells (p<0.05).

Conclusion
Id2, Id3, and Bmp4 transcripts are elevated in normal, undamaged BPs but decreased after damage. Addition of BMP4 or overexpression of Id3 after damage antagonizes Atoh1 expression and/or hair cell regeneration. These results suggest that BMPs and IDs may maintain supporting cell quiescence in undamaged tissue, and their downregulation upon hair cell loss may allow Atoh1 to drive hair cell differentiation. Future studies may focus on the manipulation of these additional factors to increase the efficiency of stimulating hair cell regeneration in the mammalian inner ear.
Capacity of consonant recognition in normal hearing and cochlear implant users

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Background
The present study sought to measure listeners’ capacity for processing multiple speech sounds, using information theory (Miller; Amer. Psychol., 1953). Normal hearing (NH) listeners are generally able to recognize more than one speech sound when two are presented, albeit with limited accuracy. We hypothesized that, if cochlear implant (CI) users’ disadvantages are limited to the periphery, their capacity (a relative assessment between the single-presentation and concurrent-presentation conditions) would be similar, although their absolute performance might be poorer than the NH group. A contrasting hypothesis was raised that, due to the severely impoverished spectral resolution in the periphery, CI users would fail to segregate concurrent sounds, resulting in drastically lower capacity.

Methods
Consonant identification (16 consonants in /aCa/ format) was tested in eight NH listeners and four bilateral CI users, who were considered good users. After establishing baseline performance with one target sound, subjects were presented with two targets and asked to identify both. The two stimuli were presented monaurally or dichotically, and with a time delay of 0, 100 or 200 milliseconds. The relative information transmitted (RIT) for the baseline condition and composite RITs for the concurrent presentations were calculated, as defined by the sum of RITs for two target consonants, which would be 2.0 if both targets were always correctly identified.

Results
For the NH group, the mean baseline RIT was 0.93. The composite RIT for the condition of 0-ms delay was 1.22, a 32% increase from the single RIT, for the monaural condition, and 1.51, a 62% increase from the single RIT, for the dichotic condition. These results were consistent with preliminary findings reported earlier (Kwon, 2001; ASA meeting abstract 1pSC27). The composite RITs improved as the inter-stimulus delay increased, indicating listener’s ability to utilize the temporal separation between stimuli. For the CI group, the mean baseline RIT was 0.75, slightly lower than the NH counterpart. The composite RIT for 0-ms delay was 0.83 and 0.88, a 10% and 17% increase from the single RIT, for monaural and dichotic conditions, respectively. Furthermore, they did not demonstrate an improvement of RIT with an interstimulus delay of 100-ms.

Conclusion
CI users’ ability to recognize multiple sounds presented concurrently is marginal, even for good CI users. They do not appear to take advantage of dichotic presentations or temporal delays between stimuli, as much as shown in the NH group.

Vocoder Simulations of Focused Cochlear Stimulation with Limited Dynamic Range and Discriminable Steps

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Background
There is recent interest in focused stimulation of the cochlea via modalities such as tripolar electrical and infrared neural stimulation (INS). Previous experiments have examined the effect of excitation spread on speech recognition, but these were limited to models representing monopolar and bipolar stimulation, with attenuations of -2dB/mm and -8dB/mm, respectively. Focused stimulation can provide narrow-extent excitation of spiral ganglion cells with spatial attenuation of -10 to -17 dB/mm. Additionally, stimulation dynamic range and minimum discriminable steps can vary among patients, and many have looked at recognition scores as a function of these parameters. There is a void in the literature, however, regarding potential benefits of focused stimulation in cases of low signal to noise ratios (SNR), limited dynamic range, and/or few discriminable steps. The purpose of this work was to employ vocoder-based experiments to investigate speech recognition under the above conditions with synthesis filter slopes representative of both standard monopolar electrical and narrower extent excitation.

Methods
Vocoder simulations were used to assess the intelligibility of sentences, consonants, and vowels that were vocoded and presented to normal–hearing listeners for identification. Slopes of the vocoder synthesis filters were chosen to reflect the spatial spread of the stimulus longitudinally along the scala tympani. Comparisons were made between simulations
representing standard monopolar electrical stimulation and the more focused approaches. Intelligibility was assessed for the different filter slopes under a variety of SNR levels, dynamic ranges, and numbers of discriminable steps.

**Results**
Speech processed via vocoder simulations representing focused stimulation was found to be substantially more intelligible than speech processed via a monopolar electric vocoder simulation. In noisy conditions, differences as large as 60 percentage points were noted (Figure 1). The data from Fig. 1 also indicate that there were no significant differences between the two filter slopes for focused stimulation. Speech processed via the focused vocoder was robust to constraints on small envelope dynamic range and small number of discriminable steps within the dynamic range. High performance was maintained with a 5-dB dynamic range and 8 or more discriminable steps. Significant drops in intelligibility were noted when the number of steps fell below 8.

**Conclusion**
Focused stimulation – tripolar electrical and INS - has potential for increased performance in noise compared to monopolar stimulation. INS holds additional value because it can achieve selective activation at each channel independently, which may lead to other benefits as well, including in music perception.
The Contribution of Room Exposure and Semantic Context to Speech Intelligibility in Cochlear Implant Users

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Background
Speech is often embedded in environmental noise and is distorted in a variety of ways such as reverberation. Previous studies on speech intelligibility in reverberant listening environments on normal-hearing (NH) listeners found facilitative effect of semantic context on increased intelligibility and prior exposure to the reverberant listening environment improved intelligibility. However, there have been no studies that investigate the effect of prior room exposure and semantic context on speech intelligibility in cochlear implant (CI) users. The current study investigates the cumulative effect of reverberation, noise, and semantic context on speech intelligibility in CI users.

Methods
Virtual acoustic techniques were used to simulate two listening environments – low reverberant (LR) room (T60 = 0.1s) and high reverberant (MR) room T60 = 0.25 s). CI users were presented with Speech Perception in Noise (SPIN) test sentences in quiet and in +15 dB signal-to-noise ratio. The carrier phrase and the target word of the speech stimuli were manipulated to be either in the same room or different rooms. Stimuli in congruent condition enhanced prior room exposure by having the carrier phrase and the target word in the same room. Stimuli in incongruent condition limited the prior room exposure by having the carrier phrase and the target word in different rooms. Listeners were instructed to repeat the last word of every sentence. Speech intelligibility was measured as the proportion of final words correctly identified by the CI users.

Results
Predictable sentences had higher speech intelligibility as compared to unpredictable sentences (context effect) in both congruent and incongruent conditions. This was true for both silent and noisy presentation conditions. The context effect was higher in LR listening environment as compared to MR listening environment. Increased intelligibility due to prior exposure (adaptation effect) was higher predictable sentences compared to unpredictable sentences in both congruent and incongruent conditions.

Conclusion
Semantic context of speech has a facilitative effect on speech intelligibility for CI users. This effect was larger in LR condition as compared to MR condition. Also, prior exposure to the reverberant listening environment helps in improving the speech intelligibility of CI users. Overall, semantic effect has a greater effect on speech intelligibility as compared to prior exposure to the reverberant environment.
Preliminary Estimates of F2 Formant Discrimination Thresholds in Cochlear Implant Users
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Background
Formant discrimination thresholds (FDTs) reflect the smallest change in formant center frequency a listener can perceive. As a measure of spectral acuity obtained with a speech-like stimulus, FDTs may provide useful insights concerning the factors that limit vowel perception by cochlear implant (CI) users. FDTs have been studied extensively in young normal-hearing (YNH) listeners, but have not yet been studied systematically in CI users.

Methods
Second formant (F2) FDTs were obtained from three CI users and three YNH listeners. Procedures and stimuli were modeled after the low-uncertainty procedures used by Kewley-Port and Watson (1994, JASA 95, 485-96) but with shorter durations of testing and an expanded range of F2 values. Stimuli were Klatt-synthesized versions of three target vowels ("uh", "ae", "ih") with 29 F2 frequency values per vowel. FDTs were measured for both increments and decrements in F2 using an adaptive 3AFC task with feedback. Each FDT was based on threshold estimates for six consecutive 80-trial stimulus blocks.

Results
FDTs for the three YNH listeners were comparable to those reported by Kewley-Port and Watson in highly trained listeners: average FDTs were 2.4% of reference frequency in the present study compared to 1.5% in the earlier study. FDTs were successfully obtained in all three of the CI users. Two of the CI users exhibited FDTs that were relatively consistent across F2 frequency, but were about 70% larger than mean FDTs for the YNH listeners. The third CI user demonstrated more variable FDTs that approached the YNH values in one frequency region but were enlarged substantially in another region. FDTs in this subject could not be explained by place-pitch thresholds obtained in an earlier study, suggesting that spectral acuity for complex stimuli may not be directly predictable from measures of spectral acuity for simple stimuli.

Conclusion
1) Our reduced-duration protocol (about 1 hour per FDT) produces FDTs in YNH listeners that are comparable to those reported for highly-trained listeners. 2) It is feasible to measure F2 FDTs in CI users by employing an expanded range of F2 increments. 3) As expected, CI users demonstrate FDTs that are larger than corresponding FDTs in YNH listeners. Future studies will determine whether variations in FDTs among CI users can be predicted by pure tone frequency DLs and whether formant discrimination predicts vowel perception in these listeners.

Phonemic Restoration and Continuity Illusion of Speech in Users of Cochlear Implants
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Background
Understanding speech in noisy environments is more difficult for cochlear implant (CI) users than normal hearing (NH) listeners. This may partially be because NH listeners are better at using the mechanism of phonemic restoration (PR) to their benefit than CI users. PR is perceptual restoration of speech sounds obliterated by noise using top-down processes.
In NH, PR is accompanied by continuity illusion, which makes listeners perceive the speech stream to be continuous behind the noise despite inaudible parts. In the present study, we explored the hypothesis that typical CI users may not benefit from PR as much as NH listeners and the continuity illusion of interrupted speech may also work differently.

**Methods**

Dutch CI users listened to the context-rich sentences. Some sentences were temporally interrupted with silence, while in others the temporal interruptions were filled with noise at various signal to noise ratios. The participants listened to and repeated the processed stimuli. Percent correct of the repeated words was measured as the intelligibility score. Benefit from PR was measured as the increase in intelligibility score with addition of noise in the silent gaps. For a given condition, continuity illusion was measured as the percent of interrupted sentences reported to be perceived as continuous.

**Results**

In the preliminary data obtained, the participants generally performed worse in identifying the sentences with noise filled gaps than silent gaps. Only two CI users (out of five tested) showed some restoration effect at a small number of test conditions. However, in general, as the level of the filler noise increased the speech intelligibility became worse. Sentences interrupted with noise were not always reported to be continuous.

**Conclusion**

Except a small number of conditions for a subset of participants, the CI users seemed to show no systematic benefit of PR in general. In contrast with the results found with the NH listeners in the previous studies, the filler noise seemed to hinder the intelligibility of interrupted speech for the CI users. As opposed to NH listeners, CI users also did not provide a clear indication of continuity illusion induced by filler noise.

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**Factors that Predict the Perception of Brief Prosodic Features by Postlingual Cochlear Implant patients**

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**Background**

While the perception of prosody has received some research attention only a limited number of studies have reported on the abilities of Cochlear Implant (CI) listeners to perceive brief prosodic cues. In Scandinavian languages these cues can fulfill a semantic role.

**Methods**

Testing of the perception of brief prosodic features was undertaken with Danish (n=18) and Swedish cohorts (n=22) of CI listeners. A normal hearing control group from both languages was also tested. Closed set minimal pair testing of lexical stress, vowel quantity, compound word/phrase discrimination and stød. With the Swedish listeners a noise condition was also included. The results from these tests were used as the dependent variable in a regression model. The factors in the regression model also included speech reception threshold results from a speech in noise test, demographic and psychoacoustic variables. The psychoacoustic variables tested with both groups included the Difference Limens for F0, intensity, duration and F1, which were measured with manipulation of a synthetically produced phoneme [ba].

**Results**

CI participants could not identify brief prosodic cues as well as normal hearing participants and their performance decreased significantly with the addition of noise. With the Danish CI listeners the SRT from a sentence test in noise emerged as a strong predictor. In the Swedish CI listeners age and noise condition emerged as predictors. In both languages the Difference Limen for intensity emerged as a predictor of performance on the minimal pair tasks. Analysis of the error patterns observed and the stimulation delivered by a common processing strategy reveals that electrical over-representation of the lower formants may be detrimental to perception.

**Conclusion**

This study shows that CI listeners are not able to perceive brief prosodic cues in natural utterances as well as normal hearing listeners.
Effects of context on speech recognition by cochlear implants users
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Background
The process of speech perception is invariably affected by the context available to the listener. The relationship between speech recognition scores in conditions with and without contextual information was modeled by the equation: \( p_c = 1 - (1 - p_i)^k \), where \( p_c \) and \( p_i \) were the scores with and without context, respectively (Boothroyd and Nittrouer, 1988). The exponent \( k \) ("k-factor") indicates the degree to which a listener utilizes contextual information. It is unclear how much contextual information is utilized in speech recognition by successful cochlear implant (CI) users. We hypothesized that CI users would rely on context to a greater degree than NH listeners to compensate for severely impoverished information from the periphery. In the present study, speech recognition scores were controlled by either adding background noise or reducing the presentation level of speech. While these two methods are considered conceptually equivalent in the calculation of articulation index, we hypothesized that they might address different implications in speech perception in CI. The noise might have a more detrimental impact because it might interfere with segmentation of the sentence into smaller linguistic units.

Methods
Ten young, NH listeners and ten “good” CI users participated in the study. 100 sentences with 4 target words were prepared and presented either with noise at several signal-to-noise ratios (SNRs) or without noise but at several reduced presentation levels. Additionally, 4 words were randomly extracted from the sentence sound files and presented as a sequence of words. Of 400 unique words, each was presented twice—one in a sentence and once in a random word string, in random order, with identical SNRs or presentation levels between presentations. The scores in the two conditions were used to calculate k-factors.

Results
While the k-factors were similar (2.3 and 2.2) for NH listeners, they were significantly different (2.9 and 2.5) for CI listeners, for the level-adjusted and noise conditions, respectively.

Conclusion
Successful CI users who perform well in daily listening tasks in the quiet condition (but with varied presentation levels) use contextual information to a greater degree than NH listeners. The well-known phenomenon of good CI users’ performance worsening sharply in noise might be attributed to the deprivation, or less use, of contextual information. Thus, more emphasis ought to be made in future studies on speech perception regarding the interference of noise with segmentation, rather than audibility per se.

Can Interrupted Speech be Used as Training Stimuli for Cochlear-Implant Users?
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Background
Normal-hearing (NH) listeners use several mechanisms that increase speech intelligibility in difficult listening environments. In phonemic restoration (PR), the listener uses context and speech redundancy to perceptually restore inaudible or masked portions of temporally interrupted speech. Previous research has shown that this benefit is greatly reduced in untrained cochlear-implant (CI) users.

CI-users commonly complain about difficulties understanding speech in noise. Several training studies have focused on improving intelligibility, although none of them have used complex stimuli like interrupted speech. We investigated if CI-users can learn to use high-level cognitive repair mechanisms more effectively to understand interrupted speech, as well as derive a PR-benefit (i.e. increase in intelligibility between sentences interrupted by silence and interrupted sentences with filler noise).

Methods
The first experiment was spread over three days with 3 sessions per day, each session comprising of 26 Dutch sentences. The NH-participants (N=30) were tested on the perception of speech interrupted by silence with or without filler noise at the start and the end of the experiment. In-between these baseline measurements, the participants were trained
on interrupted speech. Feedback was provided during these training sessions by playing back the sentence stimulus, as well as showing the sentence on the screen.

In the second experiment, with an equal design to the first, different listeners (N=24) were trained with CI-simulations of interrupted speech, by means of an 8-channel noiseband vocoder.

**Results**

First experiment: Training participants with interrupted speech without CI-simulations showed an increase in intelligibility of 9.6%. The PR-benefit was shown before training by an increase of 9.2% in correctly repeated words after the filler noise was added, and of 8.7% after training.

Second experiment: Training with CI-simulations of interrupted speech increased the performance from 17% to 37%. Consistent with previous studies, no significant PR-benefit was observed before training, and even after intensive training, such benefit was not present.

**Conclusion**

CI sound transmission degrades the quality of the bottom-up speech signal, withholding high-level cognitive repair mechanisms initially to benefit from the filler noise. Even though intensive training with interrupted speech did not revive the PR-benefit in CI-simulations, overall there was a significant improvement of intelligibility of interrupted speech. This finding indicates that listeners (and potentially CI-users) can be taught to use the top-down mechanisms more efficiently to restore interrupted speech.

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**Unilateral Adaptation of the Normal Human Angular Vestibulo-Ocular Reflex**

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**Background**

A recent study showed that the angular vestibulo-ocular reflex (VOR) can be better adaptively increased using an incremental retinal image velocity error signal compared with a conventional constant large velocity-gain demand (x2). This finding has important implications for vestibular rehabilitation that seeks to improve the VOR response after injury. However, a large portion of vestibular patients have unilateral vestibular hypofunction, and training that raises their VOR response during rotations to both the ipsi and contralateral side is not usually ideal. We sought to determine if the vestibular response to one side could selectively be increased without affecting the contralateral response.

**Methods**

We tested 9 subjects with normal vestibular function. Using the scleral search coil and head impulse techniques, we measured the active and passive VOR gain (eye velocity / head velocity) before and after unilateral incremental VOR adaptation training, consisting of self-generated (active) head impulses, which lasted ~15 mins. The head impulses consisted of rapid, horizontal head rotations with peak-amplitude 15°, peak-velocity 150°/s and peak-acceleration 3000°/s².

**Results**

The VOR gain towards the adapting side increased after training from 0.92 ± 0.18 to 1.11 ± 0.22 (+22.7 ± 20.2%) during active head impulses, and 0.91 ± 0.15 to 1.01 ± 0.17 (+11.3 ± 7.5%) during passive head impulses. During active impulses the VOR gain towards the non-adapting side also increased by ~8%, though this increase was ~70% less than to the adapting side. A similar increase did not occur during passive impulses.

**Conclusion**

This study shows that unilateral vestibular adaptation is possible in humans with a normal VOR; unilateral incremental VOR adaptation may have a role in vestibular rehabilitation. The increase in passive VOR gain after active head impulse adaptation suggests that the training effect is robust.
VOR gain and time course during head impulses in patients with bilateral vestibulopathy according to etiology.

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Background
Vestibular dysfunction may involve loss of the phasic Type I vestibular hair cells and/or of the tonic Type II hair cells. We investigated the differential effect of Type I vs. II hair cell loss on vestibulo-ocular reflex (VOR) dynamics by evaluating two groups of patients with bilateral vestibulopathy: 1) secondary to systemic aminoglycoside therapy (AG; causing mostly Type I hair cell damage) and 2) bilateral Meniere’s disease (MD; associated with Type II hair cell damage).

Methods
We recorded manually-induced head impulse tests (HITs) from 19 patients with bilateral vestibulopathy using video-oculography. Eight patients had aminoglycoside therapy (AG group) and 11 patients had bilateral Meniere’s disease (MD group). Since hair cell groups respond at different rotational frequencies, we recorded HITs with low (<200deg/s) and high (>200deg/s) head velocities. We also considered HIT gains (eye velocity/head velocity) at different time points during the head impulse to assess whether eye velocities were associated with particular head movement parameters.

Results
MD patients had higher, near-normal gains in the high and low velocity paradigms compared to the AG group (p<0.001 for both paradigms). MD patients had peak gains at the point of maximal head velocity. AG patients had a peak VOR gain (0.58 ± 0.30) at the time of maximal head acceleration in the high velocity paradigm, which was followed by an abrupt drop in gain value (referred to as a ‘bump’). Gains reached a minimum value at the time of zero head acceleration (AG: 0.10 ± 0.17). Regression analysis showed that, whereas in the MD group eye velocity linearly correlated with head velocity, in the AG eye velocity followed head acceleration.

Conclusion
These differences in the VOR-generated eye movements during the HIT could be explained by a differential hair cell loss and compensation pattern in these two groups BV patients.

Analysis of eye tracking test and optokinetic nystagmus test by original video-oculography, HI-VOG.

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Background
It is essential to use an infrared CCD camera in clinical examination of the vestibular system. Devices are currently available that can quite accurately record human eye movements, based on the principle of video-oculography (VOG). We devised an original video-oculography (HI-VOG) system using a commercialized infrared CCD camera, a personal computer and public domain software program (ImageJ) for the data analysis.

Methods
The video image from the infrared CCD camera was captured at 30 frames per second at a resolution of 640*480 pl.. For analysis of the horizontal and vertical components, the X-Y center of the pupil was calculated using the original macro. For analysis of torsional components, the whole iris pattern, which was rotated by 0.1 degree, was overlaid with the same area of the next iris pattern, and the angle at which both iris patterns showed the greatest match was calculated.

Results
For quantitative analysis, the slow phase velocity of each occurrence of nystagmus, the average value of the slow phase velocity and the visual suppression value, were analyzed automatically. Analyses of the eye tracking test, as well as the test for optokinetic nystagmus were also applied in this system.

Conclusion
Using the HI-VOG system, it was possible to inexpensively perform analysis from video images recorded with an infrared CCD camera, including eye tracking test and optokinetic nystagmus test.
Modulation of oVEMP amplitudes by lateral head tilts
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Background
Recently, the ocular vestibular evoked myogenic potentials (oVEMP) have emerged as a tool for assessment of utricular function. However, there is still a lack of unequivocal evidence in humans for either the sacculus or the utriculus being the origin of the ACS oVEMP. To overcome this problem, we sought to alter the baseline activity of utricular afferents by lateral head tilts in order to achieve a modulation of the ACS oVEMP response.

Methods
In 20 healthy subjects, oVEMP were recorded in three positions: supine, right lateral (right ear down), left lateral (left ear down). The stimulus was delivered at intensities of 100 dB nHL using a tone burst stimulus at 500 Hz, with the parameters 2/2/2 ms rise/plateau/fall time at a repetition rate of 5/s.

Results
There is a large and highly significant decrease in amplitudes in the ear-down position compared to the supine position. The derived lateral tilt modulation index (LATIMO-Index) of the ACS oVEMP in this group of subjects was \(0.71 \pm 0.27\) (mean/SDV) for the ear-down position.
There is a small not-significant increase in amplitude in the ear-up position compared to the supine position.

\begin{itemize}
\item \textbf{Table 1: oVEMP amplitudes (mean/SD) in supine vs. ear-up and ear-down lateral positions.}
\end{itemize}

\begin{table}
\begin{tabular}{ccc}
Supine & Ear-up & Ear-down \\
Mean & 3.97 & 2.51 & 4.02 \\
SD & 2.27 & 1.15 & 1.85 \\
p value & 0.000 & 0.914 & \\
\end{tabular}
\label{tab:ovemp_result}
\end{table}

Conclusion
The effects of the lateral tilts observed in our study correspond well to the effects of gravity on the utricular macula. The ear-down position has the opposite gravitational effect compared to the ear-up position, which is in line with the functional anatomy of the utricular macula. Likewise, the ear-down position markedly decreased the oVEMP amplitude, whereas the ear-up position slightly increased the oVEMP amplitude. We propose that this new parameter is a part of the ACS oVEMP response that is dependent on utricular function. Therefore, it is a promising new candidate for the specific evaluation of utricular function in humans.
Development of Accelerometer Based Device for Treatment of Benign Paroximal Positional Vertigo

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Background
Purpose: Benign Paroximal Positional Vertigo (BPPV) is well treated by proper diagnosis and otolith reposition maneuver. On the other hand, high recurrence rate of 15 to 50 % causing timely, economic burdens as well as psychosocial burden in daily life. We introduce current work for development of accelerometer-based medical device to assist BPPV treatment.

Methods
Method: We developed head-fixed lightweight accelerometer sensor which transmit positional information of head to the device by using short-range wireless communication. The device has the application program with diagnostic and treatment algorithms for BPPV. It was designed to support patient to assist self-reposition or support physician to apply proper procedure of reposition maneuver.

Results
Result: We developed device under the protocol of Inha Medical Device Development Center, and confirmed proper function of sensors, device-sensor interworking and interface of application.

Conclusion
Conclusion: We expect accelerometer-based medical device contribute the patient and physician for proper management of BPPV.
Efficacy of vibrotactile vestibular neurofeedback training based on objective body sway measures in everyday life conditions

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Background
Vestibular rehabilitation strategies mostly require a long-lasting training, which is finally not often successful. An individualized neurofeedback training which is based on a body sway analysis in everyday life conditions seems to be a more promising approach.

Hence, the present study was aimed at investigating the efficacy of an individualized vibrotactile neurofeedback training for vestibular rehabilitation.

Methods
One hundred five patients who experience one of the following balance disorders for more than 12 months were included in the study: canal paresis, otolith disorder, removal of an acoustic neuroma, microvascular compression syndrome, Parkinson’s disease, and presbyvertigo. Vibrotactile neurofeedback training was performed in a double-blinded placebo controlled study design daily (15 min) over 2 weeks with the Vertiguard® system in those 6 tasks of the Standard Balance Deficit Test with the most prominent deviations from the normative values.

Trunk and ankle sway, dizziness handicap inventory, and vestibular symptom score were measured in the verum and placebo group before the training, on the last training day and 3 months later.

Results
A significant reduction in trunk and ankle sway as well as in the subjective symptom scores were observed in the verum group. Such an effect could not be found in any of the outcome parameters of the placebo group.

Conclusion
The vibrotactile neurofeedback training applied in the present study is a highly efficient method for the reduction of body sway in different balance disorders. Since the rehabilitation program is easy to perform, not exhausting, and time saving, elderly patients and those with serious, long-lasting balance problems also can participate successfully.

An Enhanced Safe Direct Current Stimulation System to Improve Vestibular Prosthesis Performance

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Background
In previous acute vestibular prosthetic stimulation experiments in chinchillas, we showed that the range of head velocities accurately encoded by a pulse-frequency-based vestibular prosthesis can be nearly doubled by suppressing spontaneous vestibular afferent activity in the implanted labyrinth using low-amplitude anodic direct current (DC). To overcome the safety concerns associated with direct current stimulation, we developed a safe DC stimulator prototype (SDCS1). This device delivers alternating current across the metal-saline interface for each of two metal electrodes but effectively delivers DC to the tissue using synchronously switched valves arranged in a bridge circuit. The fidelity of the SDCS1 output was degraded by periodic interruptions in current flow due to non-ideal behavior of the mechanical valves used in that device. In the present study, we sought to develop a next generation safe DC stimulation system, SDCS2, which solves the problems with the current flow interruptions of SDCS1 and miniaturizes the system sufficiently to enable chronic experiments of this technology in rodents.

Methods
Conceptually, the SDCS2 uses two SDCS1 subsystems attached in a dual H bridge to prevent current flow interruptions by sequentially alternating control from one SDCS1 subsystem to the other. At any time, one SDCS1 controls stimulation current flow, while the second (non-stimulating) SDCS1 switches its valves. We tested the ability of this two-system design to eliminate current flow interruptions by constructing a bench prototype using laboratory tubing, two isolated
current sources and manually operated clamps in place of the mechanical valves. The current sources alternated control in 20s intervals to allow time to manually operate the valves. Subsequently, we designed a miniaturized 3x3x1.5 cm3 SDCS2 using layered construction compatible with MEMS processes.

Results
Experiments with the bench prototype SDCS2 show that this enhanced design completely eliminates the DC flow interruptions seen with a single SDCS1. The first miniaturized prototype of the SDCS2 was built using a 3D printer and NITINOL “muscle” wires to actuate the valves. We verified that the activated valves can be operated at 0.5 Hz as expected. When a valve is closed, its impedance is over 20x greater than when it is open.

Conclusion
These results indicate that the SDCS2 should be effective at chronically delivering uninterrupted yet biologically safe DC current. Using the SDCS2, we anticipate that we can markedly expand the range head velocities that can be encoded by a vestibular prosthesis.

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Electrocochleography and ABR variability are Associated with Postsurgical Hearing Outcomes in Superior Semicircular Canal Dehiscence
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Background
Recent findings in patients with superior canal dehiscence (SCD) have shown an elevated ratio of summating potential (SP) to action potential (AP) as measured by ECochG. Changes in this ratio can be seen during surgical intervention. The objective of this study was to evaluate the utility of intraoperative ECochG and ABR as predictive tools for post-operative hearing outcomes following surgical plugging via middle cranial fossa approach for superior semicircular canal dehiscence syndrome (SCDS).

Methods
This was a review of 23 cases (22 patients) in which good, reproducible intraoperative ECochG recordings were obtained during surgery. Diagnosis of SCDS was based on history, physical examination, vestibular function testing, and computed tomography imaging. Simultaneous intraoperative ECochG and ABR were performed. Pure tone audiometry was performed pre-operatively and at least 1 month post-operatively. Changes in SP/AP ratio, SP amplitude and ABR wave I latency were compared to changes in pure tone average (PTA) before and after surgery.

Results
SP amplitude and SP/AP ratios were reliably obtained in the affected ears in these cases. Reproducible ABR recordings were found in 19 affected ears. Mean pre-plugging SP/AP ratio was 0.64 (SD 0.33) and decreased to 0.48 (SD 0.40) immediately after plugging (p=0.015). Mean pre-plugging SP amplitude was 0.25 µV (SD 0.22µV) and decreased after surgery to 0.17 µV (SD 0.22µV) (p=0.033). Mean ABR wave I latency was 2.54 ms (SD 0.26 ms) and was unchanged after surgery (mean 2.48 ms (SD 0.32ms), p=0.48). Before surgery, mean air-conduction PTA was 17.6 dB (SD 8.9dB) and increased to 20.9 dB (SD 17.3 dB) at least 1 month post-operatively. Pre-operative bone-conduction PTA was 9.1 dB (SD 9.8dB) before plugging and 16.5 dB (SD 21.2dB) at least 1 month after surgery. A significant correlation was found between ABR wave I variability and change in postoperative PTA (p=0.03). Intraoperative change in SP amplitude was also associated with worse PTA (p=0.02).

Conclusion
This study confirmed the presence of an elevated SP/AP ratio in ears with SCD. Intraoperative ECochG and ABR variability may predict hearing outcomes in patients with SCDS.
Reducing sound exposure during Ocular Vestibular Evoked Myogenic Potential Testing for Superior Semicircular Canal Dehiscence Syndrome

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Background
Ocular vestibular evoked myogenic potentials (oVEMP) testing in response to air-conducted sound (ACS) has demonstrated excellent sensitivity and specificity for the diagnosis of superior semicircular canal dehiscence syndrome (SCDS). However, some patients experience oscillopsia while undergoing the test. Purpose: To develop an oVEMP testing protocol that may reduce the discomfort of the test and still serve for the diagnosis of SCDS.

Methods
Subjects: Five (ears= 10) healthy volunteers without history of neurotological complaints and ten (affected ears= 10) patients with a diagnosis of SCDS based on clinical presentation, audiometry and CT imaging.
VEMP recordings: OVEMPs where recorded in response to ACS (clicks and 500 Hz tone-bursts) delivered at the standard repetition rate of 5 pulses per second (pps) during 20s. Additional experimental recordings consisted of ACS delivered at a repetition rate of 2pps during 20s and 10s. Receiver operator characteristic analysis, evaluated the effectiveness of these different oVEMP protocols.

Results
OVEMP amplitudes from the three different protocols showed excellent (>90%) sensitivity and specificity for the segregation of patients with SCDS from controls. For controls, the experimental recordings with minimized sound exposure showed oVEMP amplitudes that were reduced (absent in 3 cases) relative to the standard oVEMP protocol. In contrast, in the SCD group oVEMP amplitudes showed no difference between protocols [F (2,27)= 0.04, p = 0.98].

Conclusion
Reducing the amount of sound delivered during oVEMP testing results in a more comfortable and tolerable test without compromising its effectiveness for the diagnosis of SCDS.
Body sway analysis in patients with Superior Canal Dehiscence
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Background
The principal aim of this study was to determine if patients with superior canal dehiscence syndrome (SCDS) have increased body sway, especially when exposed to loud sound.

Methods
Subjects: Groups were patients with SCDS (N=5) and healthy (non-age-matched) controls (N=4). The diagnosis of SCDS was based on clinical presentation, audiogram, vestibular evoked myogenic potentials, and high-resolution CT imaging.

Methods: Small, wireless motion sensors, positioned at the hip and ankle, measured angular body sway at 100Hz in the anteroposterior (AP) and mediolateral (ML) directions. Subjects stood with eyes open (EO) or eyes closed (EC) on firm support, with and without exposure to air conducted sound (ACS, 105 dB nHL clicks at 5Hz), during 20s trials. Center of mass excursions in AP and ML planes were analyzed.

Results
ML and AP sway was increased in the SCDS group for both conditions, and overall sway tended to align with the affected superior canal (SC). However, upon sound stimulation, SCDS patients showed a surprising decrease in sway amplitudes, especially in the ML direction during the EC trials.

Conclusion
SCDS patients have greater body sway amplitudes than controls, and sway direction corresponds to the affected SC. Loud clicks surprisingly decreased the amount of body sway excursion in this group of unselected SCDS patients. Future work will examine whether sound increases body sway in SCDS patients with sound-induced vertigo.
Dynamic visual acuity for screening patients with vestibular disorders
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Background
Several versions of dynamic visual acuity tests have been developed for use in testing patients with vestibular disorders. Many of these tests involve active head movements, which could allow the patient to perform the test with predictive saccades. We have developed a new test that passively moves the patient vertically while viewing a simple image on a computer screen.

Methods
Patients with known vestibular disorders and normals sat in a custom-designed chair that oscillates vertically. Each subject performed two trials with the chair motionless and two trials with the chair oscillating at 2 Hz. During each trial, the subjects identified the orientations of a series of Landholdt-C optotypes presented briefly (75 ms or 500 ms) on a computer screen positioned 4 m away. A technician recorded each answer with a numeric keypad, which then prompted the showing of the next optotype. The C varied in orientation and size based on an algorithm that accounts for the number of correct and incorrect answers.

Results
Preliminary data show significant differences between patients and normals in the difference score between stationary and moving conditions.

Conclusion
This test may be useful for clinical screening, at remote locations such as future landing sites for astronauts after space missions, and in future epidemiologic tests.

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Gaze Stabilization Test Asymmetry Score as an indicator of previously reported head injury in a cohort of collegiate football players.
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Background
The awareness and incidence of TBI due to contact sports is rising. Sport-related concussion is a complex pathophysiological process that can have lasting effects on brain function, balance, and behavior. Thus, there is a critical need to provide valid, reliable injury assessment measures to minimize the risk of recurrent injuries. It is therefore strongly recommended that a multi-faceted assessment protocol be used in the identification of concussion and the inclusion of more objective, sensitive measures are warranted. Vestibular function is currently not included in concussion management. Vestibular dysfunction may lead to decreased visual acuity with head movements, which may impede athletic performance and result in further injury. The Gaze Stabilization Test (GST) determines the maximum rightward and leftward head velocities in degrees/second that the athlete can perform and correctly identify a target on a computer screen. An asymmetry score denotes differences in the average head velocity between rightward and leftward head movements expressed as a percentage. The purpose of this investigation was to determine differences in the gaze stabilization test for collegiate football players reporting a previous history of head injury.

Methods
Forty collegiate football players, aged 18-22 years, from the University of Nebraska-Lincoln (14 with previously reported concussion) completed GST in the yaw plane. GST velocity scores for leftward and rightward head movements were measured and compared to participants’ report of previous head injury and compared to a more widely accepted postural control measure for concussion management - NeuroCom® Stability Evaluation Test (SET).

Results
Our findings revealed that athletes with a previous history of head injury had a larger asymmetry score (M = 11.94, SD = 8.97) than those without head injury (M = 4.91, SD = 4.77; t20.94=-2.863, p=0.009). This finding suggests that those with
previous history of concussion may have asymmetrical VOR performance that results in reduced gaze stability. Moreover, the GST asymmetry score was found to be the superior measure for identifying previous history of head injury when compared to the Stability Evaluation Test (SET). Receiver Operator Characteristic Curves (ROC) identified an area under the curve (AUC) of 0.76 for GST asymmetry suggesting good performance and AUC of 0.50 (ROC curve line closest to chance performance line) for SET indicating poor test performance.

**Conclusion**

Conclusions: Overall, these preliminary data provide evidence to support the use of the GST in concussion identification and management. This work may shift the traditional paradigm for concussion assessment to include objective vestibular measures.
Gait Characteristics of Patients with Vestibular Hypofunction
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Background
One of the most important functions of vestibular system is to maintain postural stability during head movements. Patients with vestibular hypofunction often complain of gait instability. Dynamic gait index (DGI) is a measurement tool frequently used to evaluate functional stability of gait for individuals with vestibular deficits. The purposes of this study were to: 1) Analyze the gait characteristics of patients with vestibular hypofunction during walking with active head movements, 2) Compare the differences of gait characteristics between healthy adults and patients with vestibular hypofunction.

Methods
Nine individuals with vestibular hypofunction (experimental group) and 10 healthy subjects (control group) performed DGI (items 1, 3, 4). Gait characteristics were captured by markers of the Vicon motion analysis system (Vicon Oxford, UK). Data was analyzed by NEXUS 1.7.1. Matlab 2010 was used to calculate gait parameters including step length, cadence, velocity, path deviation angles, and center of mass (COM) deviation for forward walking in the experimental and control groups.

Results
The step length and velocity decreased in the experimental group, especially during walking with horizontal (p=0.037) and vertical head movements (p=0.048). The experimental group showed larger mean path deviation angle (p=0.013). The experimental group demonstrated lesser head rotation/flexion-extension angles and trunk rotation during walking (p=0.026). The medio-lateral deviation ratio of COM of experimental group is greater than control group (p<0.01).

Conclusion
The vestibular hypofunction patients walk with lower efficiency than normal subjects. The differences were more pronounced during walking with vertical head rotation at 90 and 120 degrees/second, and horizontal head movements at 120 degrees/second. The path deviation angle of walking as well as medio-lateral deviation ratio of COM were the most sensitive parameters to show the differences of gait and balance performances between patients and normal subjects.

Rotary Vestibulo-Ocular Reflex Suppression and Slow Phase Velocity Differences in Autism
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Background
Vestibular processing and oculomotor control deficits are commonly reported in children with autism spectrum disorder (ASD), yet only a few studies have investigated vestibulo-ocular reflexes (VOR). Healthy and lesioned VORs have been
well studied in humans as well as animal models and serve as the optimal focus for studying the neural mechanisms of vestibular and oculomotor deficits in ASD. The presence of a functioning VOR occurs early in development so deficits may be identifiable prior to onset of core social and communication ASD symptoms.

**Methods**

Typically developing (TD) children and children with ASD ages 6-12 participated in velocity step tests and sinusoidal harmonic acceleration (SHA) tests in three conditions: (1) Light (2) Dark and (3) Visual Suppression - dark except for a single LED visible. All tests conducted using a NeuroKinetics VOG system and rotary chair customized by the authors for pediatric use.

**Results**

Preliminary data indicate differences in rotational VOR (rVOR) between groups during visual suppression conditions. First, phase lag was reduced during suppression but not dark SHA tests at 0.05 Hz and 0.10 Hz in ASD compared to TD children. Second, the children with ASD had a longer time constant during velocity step tests with suppression relative to both the light and dark conditions, compared with TD children. Finally, in the dark the ASD group exhibited greater occurrence of episodes where slow phase is followed by a failure to generate a quick phase prior to subsequent nystagmus beats.

**Conclusion**

Our results suggest that children with ASD respond differently to visual stimuli during continuous and sinusoidal rotation and exhibit abnormal compensatory eye movements in the absence of visual stimuli compared with TD children. These novel findings will serve to improve our understanding of VOR function in ASD. Furthermore, this information will help to shed light on three important issues in ASD including: (1) identification of neurobiological loci for sensory motor processing differences in ASD, (2) provision of a potential behavioral marker for study as an early identification of risk for ASD and (3) improved clinical assessment, patient-treatment matching and treatment outcome measures for sensory processing deficits in ASD. Future studies will aim to evaluate the specificity of these findings to ASD as well as potential perceptual differences resulting from rVOR deficits in ASD such as dynamic visual acuity and subjective visual vertical/horizontal.
Functional Brain Imaging of Multi-Sensory Vestibular Processing during Computerized Dynamic Posturography using Near-Infrared Spectroscopy

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Background

Functional near-infrared spectroscopy (fNIRS) is a non-invasive brain imaging method that uses light to determine regional changes in cerebral cortex blood flow in the cortex. In this study, fNIRS was used to investigate how the brain processes information from multiple sensory modalities during computerized dynamic posturography.

Methods

Subjects included fifteen healthy volunteers (9M/6F; ages 28 +/- 9 yrs). FNIRS testing was performed during sensory organization test (SOT) conditions I, II, IV, and V. Due to the nature of the functional NIRS paradigm (as is done in fMRI and other functional testing paradigms), four pairs of two SOT conditions were created. Each pair consisted of a baseline condition-test condition-baseline condition comparison where the condition had one fewer sensory system (vision or proprioception) than the baseline. A bilateral fNIRS probe was used to examine cortical brain activation from the frontal, temporal, and parietal regions.

Results

The figure shows the reconstructed image of brain activation during each four of the pairwise comparisons. Conditions that forced vestibular reliance by dropping out both vision and proprioceptive information (Figure 1A and Figure 1D) caused significant bilateral temporo-parietal activations (specifically in the STG and SMG). Conditions that only dropped out either vision or proprioception (Figure 1B and Figure 1C) caused significantly less STG/SMG activation. Thus we found that there was bilateral activation in the temporal-parietal area including the superior temporal gyrus (STG) and supramarginal gyrus (SMG) when both vision and proprioceptive information were degraded thereby forcing reliance on primarily vestibular information in the control of balance.

Figure Legend. Estimated spatial maps (T-test) of the oxy-hemoglobin data collected using functional near-infrared spectroscopy (fNIRS) to detect the change in brain activity during the test condition compared to the baseline conditions.

Conclusion

Our result is consistent with previous reports of the role of these regions in vestibular processing. This study demonstrates the potential utility of fNIRS in the assessment of cortical activity during standing balance tasks.
VOR Gain and Compensatory Saccades Immediately after Unilateral Deafferentation: A Prospective Longitudinal Study
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Background
The time course of changes in vestibular neurophysiologic function that occur immediately following unilateral deafferentation has not been well characterized in humans. We evaluated ipsilesional and contralesional vestibulo-ocular reflex (VOR) gain and metrics of compensatory saccade pre-operatively and for six consecutive days following vestibular schwannoma resection.

Methods
Vestibular schwannoma resection occurred via retrosigmoid approach at a tertiary academic medical center. Bedside video-oculography with a portable lightweight goggle system (EyeSeeCam) recorded eye and head rotations at a 250 Hz frame rate. Pre-operatively and then for six consecutive post-operative days (POD), we measured the horizontal VOR during head impulse testing and the latency of compensatory saccades.

Results
To date, we have evaluated 2 out of a projected 10 patients. On POD 1, mean ipsilesional horizontal VOR gain was 0.11 ± .008 (range 0.10 and 0.12). VOR gain on the contralesional side was reduced about 50% (mean 0.58 ± 0.03, range 0.55 and 0.60). Over the inpatient stay, ipsilesional gain remained low while the contralesional gain returned to baseline by POD 3 or 4. Ipsilesional head impulses elicited compensatory saccades that occurred after the head stopped moving (overt) during the early days of recovery (POD 1 – 4). Compensatory saccades that occurred during the head rotation (covert) emerged at POD 5. On POD 2, mean compensatory saccade latency was 214.6ms ± 35.0 ms, which reduced to 160.7 ± 62.5 ms by POD 5 – 6. Compensatory saccades occurred for contralesional head impulses as well.

Conclusion
Our data provide evidence of significant compensation beginning in the first days following unilateral deafferentation. This appears to occur via 2 primary means: normalization of the contralesional VOR gain, and a reduction in the latency of the compensatory saccades. We are continuing to explore the clinical and functional correlates of these changes in VOR gain and saccade metrics.

Does aging impact the response to vibrotactile feedback (VTF) during balance tasks?
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Background

Human balance relies on the integration of vision, somatosensation, and vestibular inputs. Vibrotactile feedback (VTF) applied to the torso provides an additional sensory input to augment postural control. The purpose of this study was to examine short-term and long-term learning while using VTF during conditions of reduced sensory input. We also examined the effect of age on learning during VTF.

Methods

Eighteen young (age: 24.7±2 y; 8 males) and twenty older (age: 75.4±6 y; 10 males) healthy adults participated. Each subject completed three study visits including one training visit and two experimental visits. The standing balance task required subjects to stand for 120 s during different reduced sensory feedback conditions using a Smart Equitest™ platform. During the second and third visit, three sensory conditions with VTF were tested in random order after a short re-training period. The conditions included: eyes open with darkened goggles on a fixed platform (EOD/FIX), eyes open in light on a sway-referenced platform (EO/SR), and EOD/SR. The root-mean-square (RMS) of the center of pressure (COP) in the anterior-posterior direction was calculated during different time-periods (1: 0-30 s, 2: 30-60 s, 3: 60-90 s and 4: 90-120 s). A repeated measures ANOVA was conducted to investigate the effect of age (Young/Older), and the effect of short-term learning (Periods 1-4) and long-term learning (Visit 2/3), as well as the interactions on the RMS COP.

Results

In the condition of EO/SR, there were significant main effects of Period (p = 0.012) and Group (p = 0.002, RMS greater in older adults). In the conditions of EOD/FIX and EOD/SR, there were significant main effects of Period (p < 0.001), Visit (p < 0.02, RMS reduced on last visit), and Group (p = 0.001, RMS greater in older adults). Generally, the post-hoc analyses showed that the RMS of Period 1 was significantly greater than the RMS of other periods. No significant interactions were found in the three different conditions.

Conclusion

The reduced RMS COP after period 1 and during visit 3 suggests that young and older adults are able to learn to use VTF over short and long-term periods. The lack of an interaction of Period and Visit with Group indicates that older adults are able to utilize the VTF as well as young adults.

Blast Exposure is Associated with Unilateral Vestibular Damage in US Veterans

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Background

Blast exposure and mild traumatic brain injury are significant problems for the current and recent conflicts, however, data remains limited on the effects of a blast on the vestibular system. The goal of this work was to determine if veterans with a blast exposure demonstrate impaired vestibular function.

Methods

Ten veterans (males, 41.8 ±8.5 years, 7 endorsed blast exposure) participated. Ocular counter-roll was measured during two different protocols. To assess bilateral response, subjects were tilted ±20 degrees at 0.03125 Hz (32 sec cycle) in the dark. To assess unilateral response from each ear, subjects were rotated in the dark at 400°/sec for 5 min after which there were translated 5 cm off center to the right and left for 30 sec, repeated 3 times. This allowed for unilateral stimulation of each otolith. Infrared images of both eyes were recorded continuously and analyzed using commercial software (SMI). Translation was not performed until subjects reported no sense of rotation.

Results

Examination of the tilt protocol demonstrated that bilateral ocular counter-roll was not different between groups (Controls: 0.108 ±0.021; Blast: 0.140 ±0.014). However, while unilateral values between sides did not differ in the control group (Max Response: 0.070 ±0.020; Other side: 0.067 ±0.016), they were significantly different in the group with Blast Exposure (Max Response: 0.124 ±0.013; Impaired side: 0.070 ±0.010).

Conclusion

These data indicate that veterans with blast exposure appear to have increased risk of unilateral otolith damage. In addition, this unilateral damage is masked when tested using stimuli that affects both ears. Further work is needed using a
The Effect of External Sound on Maintaining Balance

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Background
Maintaining balance and reducing postural sway is critical in the prevention of falls, particularly among the elderly and those with vestibular impairments. Although enhancing haptic and visual cues has been shown to facilitate improvements in balance, little research has examined the contribution of auditory cues as a mechanism for maintaining balance. What work has been done has focused on modulation of auditory cues to provide a feedback signal to reduce sway. The present study demonstrates how stationary auditory stimuli provide useful external reference cues to maintain balance in patients with sensory deficits affecting balance.

Methods
12 subjects with hearing or vestibular deficits stood while their center of pressure was measured for four conditions consisting of three 20-second trials each: (1) visual and auditory cues absent (2) visual cues only (3) auditory cues only and (4) visual and auditory cues present. Visual cues were provided by a high contrast abstract landscape and auditory cues were provided by four speakers at ear height emitting a broadband sound stimulus at 55 dB SPL. The speakers were placed at ear height and 50 cm in front of each subject, to each side, and behind. Pressure was measured at 100 samples/second. Sway was quantified as the 95% level on the cumulative distribution of pressure measurements for each 60-second trial condition.

Results
Sway with sound or vision added was linearly related to baseline sway. Sound reduced sway by 51% over the baseline condition. The improvement in sway with the addition of sound was linearly related to the improvement with addition of vision only; the addition of sound represented 55% of the addition of vision.

Conclusion
These results indicate that auditory input provides sensory information that is remarkably important for maintaining balance, particularly in low-light conditions. This is particularly important for patients with poor baseline balance performance. These findings indicate that auditory input is included in the rich multisensory combination of visual, proprioceptive, and visual cues responsible for maintaining balance. They also suggest that optimized auditory input could be used as a tool to improve balance in those with sensory deficits, including in elderly subjects at an elevated risk for falling. These auditory inputs could include environmental cues or improvements in hearing aids or cochlear implants.
Psychology of Dizziness
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Background
Dizziness is a complex symptom resulting from an interplay of physiological and psychological factors. One potentially relevant psychological factor is called Locus of Control (LOC). LOC is a well-established psychological concept that reflects the extent to which an individual believes that he/she is controlled by either external or internal factors. LOC has not been studied in vestibular patients with and without caloric test abnormalities. The original LOC has been extended to assess the extent to which internal control, powerful others and chance play in controlling our symptoms.

Methods
Patients who presented for electronystagmography testing filled out a standardized, validated questionnaire to establish their LOC in three domains – INTERNAL, POWERFUL OTHERS, and CHANCE. Eight weeks later the subjects were contacted to see what treatments they had used, if they had improved, and what treatment(s) they thought had helped. Using IBM SPSS statistics v19.0, analysis of variance was used to assess the relationship between LOC, improvement, and treatment. Single sample t-tests were used to determine whether or not the scores differed from published normal values.

Results
Our response rate was 80.5%. Ninety-four percent reported that their symptoms had either not changed or were better. Only 6% were worse. There were no differences in improvements rates due to treatment or diagnosis or ENG testing results. There was a significantly lower INTERNAL locus of control in our sample (p=0.042) but not for POWERFUL OTHERS (p= 0.705) or CHANCE (p=0.816). Patients who scored high in the POWERFUL OTHERS domain had a statistically significant greater chance of showing improvement (p=0.019) but neither INTERNAL nor CHANCE scores were significantly associated with improvement.

Conclusion
Patients with dizziness may feel that outside forces control their situation to a greater extent than the general population. LOC is not useful to guide treatment for dizzy patients.
ATP response of P2X2 receptor deafness mutation and association with prestin activity

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Background
ATP is an important extracellular signaling molecule and can influence cellular function in many aspects through the activation of purinergic (P2) receptors. P2 receptors have ATP-gated ionotropic (P2X) and G protein-coupled metabotropic (P2Y) subgroups. P2X2 is a predominant isoform and extensively expressed in the cochlea, including hair cells and supporting cells. Recently, it has been found that P2X2 mutation can induce human nonsyndromic hearing loss DFNA41. However, the deafness mechanisms remain unclear. We previously demonstrated that ATP can activate P2X receptors to mediate outer hair cell (OHC) electromotility and K+-sinking and recycling in the cochlea. In this study, we investigated whether P2X2 deficiency can affect ATP activation and prestin activity.

Methods
Prestin and P2X2 wild-type (WT) and V60L mutant were cloned and transfected into HEK 293 cells. ATP-evoked responses and prestin electromotility associated nonlinear capacitance (NLC) were recorded by patch clamp. We also used P2X2 knockout (KO) mice to assess whether P2X2 deficiency can diminish OHC response to ATP.

Results
Both P2X2 WT and V60L mutant targeted to plasma membrane. However, P2X2 V60L deafness mutation diminished ATP responses. The ATP-evoked inward current was eliminated in P2X2 V60L mutant transfected cells. We also found that ATP lacked the effect on prestin activity in prestin-transfected cells. However, the ATP effect was restored after co-transfection with P2X2 receptors. NLC was right-shift as we previously reported in native OHCs to ATP stimulation. P2X2 deficiency also eliminated the effect of ATP on prestin activity. OHCs in P2X2 KO mice retained NLC but lacked the response to ATP.

Conclusion
These data demonstrate that P2X2 V60L deafness mutation diminishes ATP responses. These studies also indicate that P2X2 deficiency compromises OHC electromotility regulation, which may result in over-activity of active cochlear mechanics under noise stress to induce hearing loss.
Tumor necrosis factor-alpha-mutant mice exhibit high frequency hearing loss

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Background
Exogenous tumor necrosis factor-alpha (TNFα) modulates hair cell death by altering the expression of apoptosis-related genes in response to noxious stimuli. However, little is known on the function of TNFα in normal hair cell physiology. In this study, we investigated cochlear morphology and auditory function of TNFα-deficient mice.

Methods
Auditory brainstem responses and distortion product otoacoustic emissions (DPOAE) to measure hearing function - Plastic sections to determine cochlear morphology and anatomical structure - Myosin VII and DAB staining on surface preparations of the cochlear epithelium for hair cell counts - Immunolabeling with anti-CtBP2 to determine synaptic ribbon counts - Scanning electron microscopy to analyze stereocilia morphology - Light microscopic inspection of auditory ossicles.

Results
At one month of age TNFα-mutant mice exhibited significant hearing loss, especially at higher frequencies, which was more profound in TNFα−/− mice compared to TNFα+/− and wild-type (WT) mice. Hearing loss did not progress any further from 1 to 4 months of age. DPOAE measurements (24 kHz) indicated outer hair cell dysfunction in TNFα−/− mice, but surface preparations showed no outer hair cell loss. However, stereocilia were lost in the basal turn and had collapsed in the middle turn of TNFα−/− mice. The synaptic ribbon counts of TNFα−/− and WT mice at four months of age were similar. There was also no difference in the morphology of the organ of Corti, lateral wall, and spiral ganglion cells in TNFα−/− mice compared to WT mice, nor were there differences in the morphology of the ossicles.

Conclusion
TNFα−/− mice show loss or malformation of stereocilia of outer hair cells. The otherwise normal cochlear and ossicular morphology and normal synaptic ribbon counts support the notion that stereociliary dysfunction is a major contributor to the observed threshold shifts.

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Mice deficient in the \( H^+\)-ATPase a4 subunit have severe hearing impairment associated with enlarged endolymphatic compartments within the inner ear

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**Background**

Mutations in the \( ATP6V0A4 \) gene lead to autosomal recessive distal renal tubular acidosis in patients who often show sensorineural hearing impairment. A first \( Atp6v0a4 \) knockout mouse model that recapitulates the loss of \( H^+\)-ATPase function seen in humans has been generated and recently published (Norgett et al., 2012). Here we present the first detailed description of the structure and function of the auditory system in these \( Atp6v0a4^- \) mice.

**Methods**

Auditory brainstem responses (ABR) were recorded in mice at 14 and 20 days and endocochlear potentials were measured in mice at 19-21 days.

Gross morphology of the inner ears at P20 was examined after clearing with glycerol and paint-filling was used for inner ears at P0 and E16.5.

for histological examination samples were sectioned and stained with Haematoxylin & Eosin (HE). We selected sections for immunohistochemistry to test different markers.

We examined inner ears at P20 by scanning electron microscopy (SEM) to check the integrity of hair cell bundles in mutants and controls.

**Results**

Measurements of the auditory brainstem response (ABR) showed significantly elevated thresholds in mutant mice at 14 and 20 days old, which indicate a severe hearing impairment. In addition, analysis of paint-filled ears and sections from E16.5 embryos revealed a severe expansion of cochlear and endolymphatic ducts in \( Atp6v0a4^- \) mice.

A regulatory link between \( Atp6v0a4, Foxi1 \) and \( Pendrin \) has been reported and we find that the endolymphatic sac of \( Atp6v0a4^- \) mice expresses both Foxi1 and Pds, which suggests a downstream position of \( Atp6v0a4 \).

These mutants also showed a lack of endocochlear potential which suggests a defect in function of the stria vascularis on the lateral wall of the cochlear duct. However the main \( K^+ \) channels involved in the generation of endocochlear potential, \( Kcnj10 \) and \( Kcnq1 \), are strongly expressed in \( Atp6v0a4^- \) mice.

**Conclusion**

Our results lead to a better understanding of the role of \( H^+\)-ATPase a4 subunit in hearing function.

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**Cochlear enlargement in mice lacking Slc26a4 is a consequence of disrupted NaCl absorption**

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**Background**

Enlargement of the vestibular aqueduct (EVA), commonly found in children with sensorineural hearing loss, is frequently associated with mutations of \( SLC26A4 \) which codes for pendrin. A mouse model of EVA, \( Slc26a4^- \), develops an abnormal enlargement of the membranous labyrinth, and fails to develop hearing. The goal of this study was to determine the ionic composition of endolymph at the time at which the enlargement develops.

**Methods**

\( K^+ \) and \( Na^+ \) concentrations in endolymph of \( Slc26a4^-/^- \) and \( Slc26a4^+/- \) mice were measured with double-barrel ion-selective electrodes. The \( K^+ \) concentration and transepithelial voltage were measured from embryonic day (E) 14.5 onward and the \( Na^+ \) concentration was measured between E16.5 and postnatal day (P) 0. Expression of mRNA and protein of genes associated with \( K^+ \) and \( Na^+ \) transport in the inner ear were examined by quantitative RT-PCR and/or immunohistochemistry.

**Results**

Between E14.5 and E17.5, the \( K^+ \) concentration in cochlear endolymph remained low at ~10 mM and began to rise at E19.5. The rise of the endolymphatic \( K^+ \) concentration was delayed in the cochlea and the utricle of \( Slc26a4^- \) mice.
Between E14.5 and P8, the cochlear expression of Kcnq1, Slc12a2 and Atp1a1 increased by ~4, ~8 and ~10 fold, respectively. Expression was similar in Slc26a4+/− and Slc26a4−/− mice. A surge in the mRNA expression of Kcnq1 and Slc12a2 was observed at the time of birth. The onset of Kcnq1 and Slc12a2 protein expression in the inner ear was observed at E19.5 in both Slc26a4+/− and Slc26a4−/− mice. At E16.5, cochlear Na+ concentration in endolymph was ~140 mM, which dropped drastically at P0. The drop in the Na+ concentration was delayed in Slc26a4−/− mice. Cochlear expression of Scnn1b and Scnn1g was increased but no difference of Scnn1a expression was found in Slc26a4+/− and Slc26a4−/− mice at ages E14.5 to P8.

Conclusion
These data demonstrate that endolymph in the phase of cochlear growth during embryonic development of the mouse inner ear is a NaCl-rich fluid, which is modified into a KCl-rich fluid just prior to birth. Further, the data suggest that endolymphatic enlargement caused by a loss of Slc26a4 is a consequence of disrupted NaCl absorption.

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Mechanical Filters in Rbl2 Null Mice: The Influence of Supernumerary Inner Ear Sensory and/or Support Cells
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Background
Numerous animal models that are characterized by the presence of supernumerary inner ear sensory and support cells have been generated by inactivating or partially inactivating genes that negatively regulate the cell cycle. This includes the Rbl2 null mouse strain that stands as a unique animal model of inner ear cell proliferation in that sensitivity to sound and distortion product otoacoustic emission (DPOAE) suppression tuning curves appear normal in spite of the presence of an extra row of inner hair cells and extra row(s) of outer hair cells in the apical half of the cochlea (Rocha-Sanchez et al., 2011; Brunette et al., 2012). In this study, the shape and properties of inner ear mechanical filters in Rbl2 null and wildtype (WT) mice were compared in an effort to determine if the presence of supernumerary sensory and/or support cells alters filter characteristics at suprathreshold levels.

Methods
To address this question, DPOAE “ratio series” were obtained in WT and Rbl2 null mice. The separation between primary stimulus frequencies was varied in an orderly manner while holding f2 constant (f1<f2), and levels of the 2f1-f2 component were plotted as a function of distortion product (DP) frequency. This manipulation produces DPOAE filter functions that represent mechanical events underlying spatial interactions between two tones in the inner ear. The level of primaries used to generate responses varied from an f1 level of 20 to 70 dB SPL, in 10 dB steps, and the level of f1 was always 10 dB higher than that of f2.

Results
The center frequency of DP level versus DP (2f1-f2) frequency curves predictably shifted in the direction of lower DP frequencies as stimulus levels increased in both WT and Rbl2 null mice; however, DP peak frequency tended to be slightly lower in Rbl2 null mice than in WT cohorts. In addition, broader mechanical filter envelopes were observed in Rbl2 null mice when compared with WT mice, and DP levels produced by Rbl2 null mice were lower than in WT mice when higher level primary tones were considered.

Conclusion
Mechanical filters of the inner ears of Rbl2 deficient mice are qualitatively similar to those representing WT mice; however, extra sensory and/or supporting cells in Rbl2−/− mice appear to influence the ratio between primary stimulus frequencies producing the peak DP, and reduce filter sharpness at moderate and higher primary stimulus levels for conditions studied thus far.
Autophagy is essential for the maintenance of cochlear outer hair cells and hearing in vivo
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Background
Auditory hair cells are responsible for the transduction of sound-evoked mechanical energy to the nerve impulses throughout the life without replacement. We here investigated the role of autophagy, an evolutionary conserved intercellular recycle pathway via lysosome, in the maintenance of homeostasis in the auditory hair cells.

Methods
1) An autophagy reporter mouse line, GFP-LC3 was examined by immunostaining for GFP using antibody. The mice has robust expression of GFP-LC3 fusion protein on the autophagosome, which is a fundamental structure of autophagy. Paraffin sections of the inner ear were investigated. 2) A mouse line disrupting autophagy specifically in the inner ear hair cells was generated by crossing Pou4f3-Cre, an hair cell-specific Cre mouse, with flox-Atg7. Atg7 is an essential gene for the formation of autophagosome and its genetic depletion results in the complete abolishment of autophagy. Tone-burst ABR at 2, 4, 8, 16, 32kHz and histological examinations were performed at 8 and 32 week old of the conditional knock-outs and their littermate controls.

Pou4f3-Cre is a kind gift from Dr. Vetter (Univ. Mississippi Medical Center). GFP-LC3 (Tg(CAG-EGFP/Map1lc3b)53Nmz: RBRC00806) and flox-Atg7 (B6.Cg-Atg7tm1.1Tch: RBRC02759), were generated by Dr. Mizushima and Dr. Komatsu (Tokyo Medical Dental Univ), respectably, and both were supplied via RIKEN BRC under their permissions.

Results
Reporter expression was observed in outer hair cells and supporting cells of GFP-LC3 but was not seen in inner hair cells. Conditional knock-outs of Atg7 suffered an accelerated hearing impairment from high frequency to the lower as their ages developed. ABR threshold was significantly elevated compared to the control. Histology showed missing outer hair cells at 32 week of age.

Conclusion
The results suggest that the autophagy is required especially for the maintenance of outer hair cells but is, interestingly, not necessarily required in inner hair cells. Further studies elucidating the precise role of autophagy in the outer hair cells, especially which protein and/or organelle is degraded by autophagy, will be investigated.
New function for Connexin26 (Cx26) in the cochlea and new mechanism for deafness induced by deficiency in Cx26

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Background
Connexin 26 (Cx26, GJB2) mutations can induce a high incidence of nonsyndromic hearing loss. The deafness-inducing mechanism, however, remains largely undetermined. Hair cells have no gap junctions and connexins are only expressed in supporting cells with Cx26 being the predominant isoform in the cochlea. Recently, we have reported that cell degeneration is not a primary cause for Cx26 deficiency-induced hearing loss (Liang et al., 2012). We have also reported that supporting cells can modify active cochlear amplification, probably by influencing outer hair cell (OHC) electromotility (Yu and Zhao, 2009). However, the molecular and cellular mechanism underlying this modification remains unclear. In this study we investigate morphological changes in cochlear supporting cells induced by Cx26 deficiency.

Methods
Cx26 deficient mice were created by loxP-Cre technique. Auditory function was measured with ABR and DPOAE techniques, and confocal and electron microscopy (EM) techniques were used to assess morphological changes in the cochlea. OHC nonlinear capacitance (NLC) was measured by patch clamp recording.

Results
Deletion of Cx26 in Deiters cells and outer pillar cells reduced DPOAE and increased ABR thresholds. However, the cochlea had normal development and the tunnel of Corti was open. Patch clamp recording showed that NLC was shifted to right in depolarization direction. In EM images Deiters cells appeared fat, with plasma membrane of adjacent cells sticking together in a way rarely observed in control animals. The microtubules in DCs and outer pillar cells also appeared to be less abundant.

Conclusion
Cx26 deficiency induces morphological changes in cochlear supporting cells, probably altering their mechanical properties. These changes might influence OHC electromotility and eventually lead to hearing loss. This study also suggests a new non-channel function of Cx26 in the cochlea.
Deiters Cells’ Attachment to the Basilar Membrane is Structurally Similar, and Mediated by the Same Adhesion Molecules, in Mice, Rats and Guinea Pigs

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Background
Deiters cells extend from the basilar membrane to the reticular lamina and, together with pillar cells and outer hair cells, structurally define the micro-architecture of the organ of Corti. We have recently described that mouse Deiters cells possess a morphologically distinct infranuclear region, and their very basal portion, known as Deiters’ feet, contact the basilar membrane in a narrow stripe extending all along the length of the organ of Corti.

Methods
Using confocal laser and electron scanning microscopy we have now continued our studies in mice and extended them to two additional species, guinea pigs and rats.

Results
We found that, just like in mice, in rats and guinea pigs the basal pole of every Deiters cell was attached to the basilar membrane within an approximately 15 µm-wide stripe adjacent to the row of outer pillar cells running all the length of the cochlear spiral. Also, as described in mice, complete detachment of Deiters cells in these other two species revealed an elliptical imprint on the top surface of the basilar membrane consisting of a smaller central structure with a very smooth surface surrounded by a rougher area, suggesting the presence of two different anchoring junctions. Immunofluorescence studies with specific antibodies suggested that the adhesion molecules in the central region would be mostly integrins, whereas the peripheral region would contain laminins and the discoidin domain receptor tyrosine kinase 1 (DDR1).

Conclusion
These results indicate that Deiters cells follow the same pattern of organization, and use the same mechanisms of adhesion to the basilar membrane, in three different species, suggesting a high degree of evolutionary conservation.

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Metabolome analysis of inner ear fluid in guinea pigs cochlea
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Background
The metabolome analysis is to analyze metabolites resulting from all cellular activity to have of the organism cyclopedically. The intracellular dynamics which are not understood by expression of mRNA and proteome analysis are found by the profile of the metabolome analysis. Various clinical conditions such as cancer and immune disease have been recently apparent by metabolome analysis with many organs and tissues. This time we examined metabolome analysis of inner ear fluid in guinea pigs cochlea using gas-chromatography/mass-spectrometry (GC/MS).

Methods
Eighteen pigmented guinea pigs (250-300g; Hartley male) with normal Preyer’s reflex were used in this study. The experimental protocol was approved by the Animal Care and Use Committee at the University of Kobe. After animals were deeply anesthetized decapitated, the temporal bones were immediately removed. Under a dissecting microscope, the round and oval windows were opened and the lymphatic fluids were taken. At the same time decapitating, plasma fluid were also taken. We extracted water-soluble metabolite from the inner ear lymphatic and plasma fluids. After freeze dry and a process of derivatizing, metabolites were measured with GC/MS-QP2010. The obtained data examined the metabolite by principal component analysis.

Results
As for the metabolite which was specific for inner ear fluid of guinea pig cochlea, it was detected 29 kinds in total. Also, the metabolite, which was significantly frequent in inner ear fluid compared to plasma, was detected totally seven kinds.

Conclusion
The present study reports for the first time that metabolome analysis of the inner ear. As results, it was detected 29 kinds in total, for the metabolite which was specific for inner ear fluid of cochlea. Also, the metabolite, which was significantly
frequent in inner ear fluid compared to plasma, was detected totally seven kinds. It seems that these results lead to clinical condition elucidation of the sensorineural hearing loss including noise-, drug- aging-induced hearing loss for a diagnostic tool and treatment therapy.

### Defining the role of integrins in the repair and regeneration of hair cells in the human vestibular system

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#### Background

Integrins are cell adhesion receptors that play important roles in physiological and pathological processes throughout the body. Integrins comprise alpha and beta subunits and the 24 αβ heterodimeric members mediate the interaction of cell–cell signalling and communication between the cell and extracellular matrix.

Lesions created by the death of a hair cell is closed by supporting cells in a manner that maintains the permeability barrier at the luminal surface of the epithelium. This controlled process relies on cell shape changes and spreading that likely involves integrins. Furthermore, the supporting cells are thought to remove these dead hair cells by a phagocytic process. Certain integrin receptors recognise apoptotic cells mediating phagocytosis of the apoptotic body.

Following hair cell loss in the vestibular system of mammals there is limited regeneration possibly via the direct phenotypic conversion of supporting cell into hair cells without an intervening mitotic event. We hypothesise that integrins play an integral role in the repair and recovery process. We are able to test this hypothesis with human vestibular tissue obtained from patients undergoing trans-labyrinthine procedures for acoustic neuromas. In this first part of the work we identify which integrins are normally expressed in human vestibular tissues.

#### Methods

qPCR was used to screen for the integrin m-RNAs specifically expressed in human vestibular explants. Upon harvesting from the patient, the tissue was immediately transferred to RNAlater for transport to the laboratory. RNA was subsequently prepared using RNAeasy (Qiagen) and reverse transcribed into cDNA. Each array was performed with cDNA from the vestibular tissue explanted from a single patient. Data was collated from individual arrays and relative expression was quantified by qPCR using the ddCT method.

Directed by the results of the qPCR analysis, the expression pattern of specific integrin proteins was localised using immunohistochemistry. Human vestibular tissue was transported in Minimal Essential Medium, subsequently fixed and labelled using antibodies with human specific reactivity.

#### Results

qPCR array demonstrated integrin subunit mRNAs ITA5, ITA6, ITA8, ITAV, ITB1 and ITB5 were expressed above background levels suggesting their presence in human vestibular tissue. Previous work in our laboratory has identified these integrins in adult mouse vestibular tissue.

#### Conclusion

The integrin profiles of human and mouse vestibular systems are similar but not identical. On-going work will continue to examine integrin protein expression and location and further elucidate their role in the repair of the sensory epithelia of the inner ear and regeneration of hair cells.
Expression changes of endothelin 1, endothelin receptor A and B in noise induced cochlea
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Background
After noise exposure, impairment of cochlear blood flow has been considered to be one of the factors of hearing loss. Although there were several evidences of reducing blood flow in the cochlea after noise exposure, the pathophysiology of blood flow change is still obscure. Endothelins are proteins that constrict blood vessels and have a key role in vascular homeostasis and works through its receptors. Recent research has shown that endothelin and its receptors were present in the cochlea and these things were regarded to have a role in regulation of cochlear blood flow. In this study, we investigated the expression changes of endothelin 1 (ET-1), endothelin receptor A (ETAR) and B (ETBR) according to auditory threshold change after noise exposure.

Methods
Mice were exposed to noise to generate transient threshold shift (TTS) and permanent threshold shift (PTS) respectively. Auditory threshold shifts were evaluated with auditory brainstem response and expression changes of ET-1, ETAR and ETBR after noise exposure were evaluated by RT-PCR.

Results
Expressions of endothelin 1, endothelin receptor A and B were changed after noise exposure in both TTS and PTS group. After noise exposure, expression of ET-1 was increased in TTS group compared to control and PTS group. Expression of ETAR was decreased just after noise exposure but increased 2 weeks after noise exposure in PTS group. Little changes of ETAR expression were noted in TTS group. Expressions of ETBR were increased in both TTS and PTS group just after noise exposure but it was sustained in PTS group after 2 week after noise exposure compared to TTS group.

Conclusion
These results suggest that expression changes of ET-1, ETAR and ETBR might be associated with hearing threshold shift and recovery after noise exposure in the cochlea.
Vascular diameter of the spiral modiolar artery is regulated by Ca\textsuperscript{2+} sparks, K\textsuperscript{+} channels and Ca\textsuperscript{2+} sensitivity
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Background
The combined action of Ca\textsuperscript{2+} sparks and Ca\textsuperscript{2+}-activated K\textsuperscript{+} (BK) channels lowers smooth muscle global Ca\textsuperscript{2+} concentration leading to relaxation of vascular smooth muscle cells and vasodilation. Loss of Ca\textsuperscript{2+} sparks or uncoupling of BK channels from Ca\textsuperscript{2+} sparks leads to increase in global Ca\textsuperscript{2+} concentration and vasoconstriction. Here, we investigated whether Ca\textsuperscript{2+} sparks and BK channels regulate smooth muscle global Ca\textsuperscript{2+} and the vascular diameter of the spiral modiolar artery (SMA) and control cochlear blood flow.

Methods
SMAs from female gerbils were cannulated with a motorized concentric pipette system, pressurized at 40 or 60 cmH\textsubscript{2}O and loaded with the fluorescent Ca\textsuperscript{2+} indicator dye, fluo4. Ca\textsuperscript{2+} sparks and Ca\textsuperscript{2+} waves were recorded in line scans and frame scans. Smooth muscle global Ca\textsuperscript{2+} and vascular diameter was recorded simultaneously using confocal microfluorometry.

Results
Ca\textsuperscript{2+} sparks and Ca\textsuperscript{2+} waves were observed in pressurized SMAs at 40 and 60 cmH\textsubscript{2}O. At 40 cmH\textsubscript{2}O, ryanodine abolished Ca\textsuperscript{2+} sparks and waves, increased the global Ca\textsuperscript{2+} and caused a vasoconstriction. In comparison, at 60 cmH\textsubscript{2}O, ryanodine-induced increase in the global Ca\textsuperscript{2+} was smaller and vasoconstriction was suppressed. The pressure-dependence of the ryanodine-induced vasoconstriction was consistent with the pressure-dependence of the Ca\textsuperscript{2+} sensitivity, which was greater at 40 than at 60 cmH\textsubscript{2}O. Treatment with 1nM endothelin-1 increased ryanodine-induced vasoconstriction at 60 cmH\textsubscript{2}O. Iberiotoxin, an inhibitor of BK channels, had no effect on either global Ca\textsuperscript{2+} or vascular diameter, although in the presence of iberiotoxin, ryanodine-induced effects were greatly diminished. Iberiotoxin increased global Ca\textsuperscript{2+} and caused a vasoconstriction in the presence of apamin, an inhibitor of Ca\textsuperscript{2+}-sensitive small-conductance K\textsuperscript{+} (SK) channels.

Conclusion
The results indicate that, in the SMA, regulation of global Ca\textsuperscript{2+} and vascular diameter by Ca\textsuperscript{2+} sparks and BK channels is modulated by SK channels and the Ca\textsuperscript{2+} sensitivity. SK channels are critical for the control of global Ca\textsuperscript{2+} whereas the Ca\textsuperscript{2+} sensitivity is critical for the control of vascular diameter.
This study was funded by NIH R01-DC04280 to PW.
The functional modification of pRb during the early development of chicken inner ear

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Background
The retinoblastoma protein pRb plays a vital role in regulating mammalian cell cycle progression, and inactivation of pRb is necessary for S phase re-entry. Further, pRb plays a crucial role in cell survival. Therefore, modulation of pRb function transiently instead of permanent deletion, is an attractive route by which proliferation and survival can be achieved for hair cell regeneration.

Methods
pRb function is regulated by different types of phosphorylation: cyclin dependent that is mediated by Ras/Raf/Mek/MAPK cascade or cyclin independent that is mediated by the physical interaction between C-Raf and pRb. Using a culture system for chicken otocysts, we investigated how two small molecules U0126 and RRD 251, inhibitor of MAPK cascade or Rb-Raf-1 interaction, affect progenitor cells proliferation and differentiation during the early inner ear development.

Results
Inhibition by both U0126 and RRD 251 in the progenitor cells led to cell cycle arrest and increased apoptosis. Further, inhibition of MAPK cascade mediated pRb phosphorylation significantly reduced the number of proliferating pro-sensory progenitors; whereas inhibition of C-Raf-mediated phosphorylation significantly reduced the number of proliferating ganglion progenitors.

Conclusion
We concluded that different pathway of pRb phosphorylation play distinct roles during the early inner ear development, with implication in modulating pRb function according their cell type specificities.
Acute Ischemic Hypoxia Affects Distinct Channels in Smooth Muscle and Endothelial Cells in the Spiral Modiolar Artery
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Background
Acute ischemia generally induces a compensatory vasodilation in the most vascular beds including the cochlear arterioles but the underlying cellular mechanism remains controversial.

Methods
Using conventional intracellular and whole-cell recording from smooth muscle (SMC) and endothelial cells (EC) in situ of dispersed from segments of guinea pigs cochlear spiral modiolar artery, we analyzed the membrane actions of acute ischemic treatment (AIT, hypercapnic hypoxia, pO2=0 mmHg, pH=6.5).

Results
The AIT for 2 to 30 min caused a reversible and partially repeatable vasodilation and hyperpolarization in the majority of cells. ACh- and 10 mM K+-induced dilation was significantly reduced, lost or reversed to a constriction during and after the AIT. Holding at -40 mV, AIT induced a reversible 10-60 pA outward current within 1-2 min in the majority of SMCs and ECs. During the AIT, the whole-cell I/V curve became stronger outwardly rectifying at voltages > -40 mV. The AIT-induced net current in isolated SMCs exhibited a voltage and time dependence and sensitivity to 1-10 mM TEA or 4-AP similar to those of voltage-gated (KV) and big conductance Ca2+-activated (BK) K+-channels. AIT also reduced gap junction coupling current in in situ SMCs. In contrast, the AIT-induced current in isolated ECs was blocked by 100 µM niflumic acid but not TEA and 4-AP; the net current showed a voltage and time dependence typical of Ca2+-activated Cl--channels (CaCC): slowly activating outwardly rectifying with a reversal potential near -30 mV.

Conclusion
We conclude that the acute ischemia in the cochlear arteriole induces hyperpolarization, dilation and loss of EDHF function by implicating activation of KV and BK, inhibition of gap junction coupling in the smooth muscle cells whereas in endothelial cells, AIT mainly activates CaCCs. Supported by NIH NIDCD DC 004716, P30 05983.

Acute Ischemia Activates Outward Rectifier Currents in Cochlear Lateral Wall Capillary Cells
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Background
Microcirculation in spiral ligament (SL) and strialvascularis (SV) of the cochlear lateral wall are critical for cochlear health, especially the highly energy-demanding auditory transduction, but the physio-pathology of the local capillary cells remains poorly understood.

Methods
Using acutely isolated capillary segments and whole-cell recording techniques, we characterized the main membrane currents and the responses to acute ischemic treatment (AIT, hypercapnic pH 6.5, pO2=0 mmHg) of the endothelial cells (EC) and pericytes (PC) in the SL and SV.

Results
We found: 1) Within physiological solutions, the PCs, but not ECs, showed a [K+]o-facilitated and Ba2+-sensitive inward rectifier current. 2) ECs, but PCs rarely, showed a significant outward rectifier current, which was inhibited by 1 mM 4-AP, 100 µM niflumic acid (NFA) but not TEA. 3) When recorded in situ of a vessel segment, both types of cells showed variable electrical coupling that can be blocked by 30 µM 18β-glycyrrheticin acid or 100 µM 2-APB. 4) AIT (3 – 24 min) caused a partially reversible and repeatable outward current at holding potential -20 mV in isolated ECs and PCs. The AIT-activated net current had I-V curves and kinetics which mostly resemble to those of Ca2+-activated Cl-channels: strong outward rectification with quick or slow time course to reach steady-state within 0.5 - 2 s voltage steps. This AIT-induced current was inhibited by niflumic acid and 1 mM TEA but not 4-AP. 5) AIT (>3 min) also inhibited gap junction coupling current.

Conclusion
We conclude that the capillary EC and PC in the SV and SL express different K+-channels, suggesting a role of PCs as the K+-sensor for blood flow regulation, and the acute ischemia activates chloride channels and BK channels likely via
elevation of cytosolic Ca²⁺ in both ECs and PCs. Supported by NIH grants DC004716 (ZGJ) and DC005983, DC 00105 (ALN).

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The involvement of ROCK pathway in the apical cell contraction in murine organ of Corti
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Background
We previously reported the morphological changes and plasticity of hair cells and supporting cells by inhibition of myosin II. In the current study, we report the contribution of ROCK and MLCK pathways to the activation of myosin II on the regulation of cell shaping using ROCK inhibitor (Y27632) and MLCK inhibitor (ML7).

Methods
By using explant cultures dissected from murine organ of corti, we compared the morphological changes of hair cells and supporting cells after the treatment with blebbistatin, Y27632 or ML7.

Results
Using Y27632, we observed morphological change less than but similar to one observed in blebbistatin treatment. After Y27632 treatment, the surface area was remarkably enlarged in supporting cells, although it was slightly but significantly enlarged in hair cells. This morphological change is recovered after two more hours of culture in the medium without Y27632. These morphological changes were not observed in ML7-treated samples.

Conclusion
Our data demonstrate that ROCK pathway is involved in the contraction of apical surface in hair cells and supporting cells through the activation of myosin II. On the contrary, the involvement of MLCK in this process seems to be minimal.

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Transfection of therapeutic genes by hydroxyapatite nanoparticle complexes in organotypic cultures of the inner ear
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Background
Gene therapy is a powerful technique for specific treatment of inner ear diseases. Hydroxyapatite nanoparticle(nHAT) is an ideal gene carrier without potential side-effects in comparison with viral vectors, such as oncogenicity, cytotoxic effect, immunogenicity, or non-specific inflammation. Therefore, nHAT has received increasing attention in gene therapy. Our previous studies demonstrated that transfection of nHAT through round window membrane in vivo was mainly ingested by dark cells in the inner ear via endocytosis. However, the transfection efficiency of nHAT to the inner ear in vitro is unknown. In current study, the explants of cochlear and vestibular organs from adult rats and neonatal rats were examined after PEI-nHAT-pEGFPC2-NT3 transfection.

Methods
Adult SASCO sprague dawley rats and post-natal day 3 rats were decapitated. The cochlear organs, macula of utricle and saccule, and crista of ampullaris were cultured in serum-free culture medium or treated with or without 1mM gentamicin. nHAT-pEGFPC2 was mixed into the culture medium with a final concentration of 500µg/ml cultured for various period. The explants were fixed with 10% formalin and then stained with TRITC-conjugated phalloidin, which preferentially labels filamentous actin in hair cell stereocilia. Specimens were also stained with Topro-3 to label nucleus. The whole mount specimens were examined under a confocal laser scanning microscope using appropriate filters to detect red fluorescein of phalloidin on the hair cells, green fluorescein of EGFP in transfected cells and blue fluorescein of Topro-3 in the nucleus.

Results
In adult organotypic culture system, EGFP were hardly detected in organs 3 hours after transfection. However, EGFP were clearly visible in vestibular hair cells, spiral ganglion neurons, epithelium of stria vascularis, and also in some damaged cochlear hair cells 30 hours after transfection. Differing from adult cultures, EGFP in neonatal cochlear culture system were not detectable 48 h after transfection. However, when cochlear organotypic cultures were pre-treated with
1mM gentamicin for 10 h, and then cultured with PEI-nHAT-pEGFPC2-NT3 material, EGFP was strongly expressed in many hair cells, spiral ganglion neurons, and epithelium of stria vascularis.

Conclusion
PEI-nHAT can easily transfect therapeutic genes into hair cells, spiral ganglion neurons and stria vascularis in the inner ear in adult rat in vitro. However, PEI-nHAT is difficult to transfect developing cochlear organs at the age of postnatal day 3 in culture condition, unless the barrier of cochlear tissue was damaged by gentamicin.

A protocol for isolation and culture of endothelial cells, pericytes, and perivascular resident macrophage-like melanocytes from the young mouse ear
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Background
The cochlear blood-labyrinth barrier tightly regulates the cochlear microenvironment for auditory function. Pericytes (PCs), perivascular resident macrophages-like melanocytes (PVM/Ms), and endothelial cells (ECs) are critical components of the blood-labyrinth barrier essential for maintaining blood-labyrinth barrier integrity. However, investigation of these cell types has been hindered by lack of a method for isolating and culturing them.

Methods
Here we describe a novel growth medium-based method for obtaining cochlear ECs, PCs, and PVM/Ms from the stria vascularis of young mice (p10 - p15). The procedure does not involve mechanical or enzymatic digestion of the sample tissue. Explants of stria vascularis, “mini-chips”, are selectively cultured in growth media.

Results
Primary cell lines are obtained in 7 - 10 days. The method is simple and reliable, and provides high quality ECs, PVMs, and PCs with a purity > 90% after two passages.

Conclusion
The newly established method is suitable for producing primary culture cells from organs and tissues of small volume and high anatomical complexity, such as the inner ear capillaries. The highly purified primary cell lines enable cell culture-based in vitro modeling of cell-cell interactions, barrier control function, and drug screening. This work was supported by National Institutes of Health grants NIH NIDCD R01-DC010844 (XS), NIH NIDCD DC R21DC1239801 (XS), and NIHP30-DC005983.
Endothelial cell, pericyte, and perivascular resident macrophage-type melanocyte interactions
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Background
The integrity of the fluid-blood barrier in the stria vascularis is critical for maintaining inner ear homeostasis, especially for sustaining the endocochlear potential, an essential driving force for hearing function. However, the mechanisms that control intra-strial fluid-blood barrier permeability are largely unknown. At the cellular level, the intra-strial fluid-blood barrier comprises cochlear microvascular endothelial cells connected to each other by tight junctions (TJs), an underlying basement membrane, and a second line of support consisting of cochlear pericytes and perivascular resident macrophage-type melanocytes.

Methods
In this study, we used a newly established primary cell culture-based in vitro model to show that endothelial cells, pericytes, and perivascular resident macrophage-type melanocytes interact to control intra-strial fluid-blood barrier permeability.

Results
When the endothelial cell monolayer is treated with pericyte— or perivascular resident macrophage-type melanocyte-conditioned media, the permeability of the endothelial cell monolayer is significantly reduced relative to an untreated endothelial cell monolayer. Further study has shown the pericytes and perivascular resident macrophage-type melanocytes to regulate TJ expression in the endothelial cell monolayer.

Conclusion
The new cell culture-based in vitro model offers a unique opportunity to obtain information on the organ-specific characteristics of the cochlear blood/tissue barrier. Our finding demonstrates the importance of signaling between pericytes, endothelial cells, and perivascular resident macrophage-type melanocytes to the integrity of the intra-strial fluid-blood barrier. This work was supported by National Institutes of Health/National Institute on Deafness and Other Communication Disorders Grants DC008888-02A1 (to X.S.), DC008888-02S1 (to X.S.), R01-DC010844 (to X.S.), and NIHP30-DC005983.
Robust Extracellular Matrix Production with Recombinant Tectorin Constructs in Mouse Cochlear Cultures

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Background
Alpha- and beta-tectorin (Tecta and Tectb) are major non-collagenous components of the tectorial membrane (TM). Tecta is a large modular glycoprotein composed of several modules; an entactin domain, a zonadhesin-like domain comprising three full and two partial vWF type D repeats, and a zona pellucida (ZP) domain. Tectb is much smaller and consists of a single ZP domain. The presence of a ZP domain in both tectorins suggests that Tecta and Tectb can form hetero- or homopolymers. It is still unclear, however, how these proteins assemble to form the TM matrix. The mechanisms of apical targeting, secretion and processing of the tectorins are also largely unexplored.

Methods
Fluorescently-tagged tectorin constructs were transiently expressed in polarised epithelial cell lines (CL4 and MDCK) to explore apical targeting, and MDCK derived cell lines stably expressing these constructs were used to study secretion and processing. Polarised epithelial spheres were generated from stable cell lines by growing cells in a mixture of collagen and Matrigel. Gene gun transfection was used to transfect the tectorin constructs into various cell types in mouse cochlear cultures. Site directed mutagenesis was used to introduce mutations that are known to cause human hereditary hearing loss.

Results
Tectorins were shown to be effectively targeted to the apical surface of CL4 cells, and the targeting of both proteins was shown to depend on the presence of a C-terminal GPI anchor sequence. With MDCK-based stable cell lines expressing Tecta or Tectb, the tectorins were detected in the media, within the cells, and on their apical cellular surface. Significant amounts of matrix were, however, not observed with either stable or transient tectorin expression in monolayer cultures of polarised epithelial cell lines. With polarised epithelial spheres there was some evidence for the production of filamentous extracellular matrix. In contrast, substantial amounts of dense extracellular matrix were observed on the apical surfaces of outgrowth zone cells when cochlear cultures were transiently transfected with either Tecta or Tectb. When expressed in hair cells, Tecta and Tectb locate to the distal tips of the hair bundle.

Conclusion
The results reveal that (i) the GPI-anchor acts as a signal for the apical targeting of Tecta and Tectb, and (ii) epithelial cells in primary cultures of the early postnatal cochlea are the best suited for studying tectorin-based extracellular matrix production.

Thinking of supporting cells as glia, lessons for studying their functions.
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There is growing evidence that supporting cells of the inner ear sensory epithelia share significant similarities with glial cells elsewhere in the nervous system, and that these cell types also have common biological roles. I will introduce some of the similarities and differences between supporting cells and glia as well as discuss our studies on the roles of supporting cell-derived trophic factors in the development, function and maintenance of the inner ear.
Müller glia as retinal stem cells
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The retina is a complex neural circuit, with sensory receptors, interneurons and projection neurons to the brain. Degeneration of one or more of these different types of neurons can lead to visual impairment and blindness. In non-mammalian vertebrates, regeneration of new neurons from supporting cells, like the Muller glia, can restore normal neuronal numbers and function, but this process is very limited in mammals. Recent work from our lab has shown the feasibility of stimulating neurogenesis from the Muller glia in mice, using reprogramming factors. This talk will review current progress and future prospects for repairing the damaged retina.

Supporting Cells as Mediators of Hair Cell Survival
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Cellular stress causes induction of both pro-apoptotic and pro-survival signals. The balance of these death and survival signals often determines whether the stressed cell lives or dies. Induction of heat shock proteins (HSPs) in response to cellular stress is a ubiquitous and highly-conserved response that can inhibit apoptosis in many systems. We recently showed that HSP induction inhibits both aminoglycoside- and cisplatin-induced hair cell death. HSP70 is required for this protective effect, and constitutive expression of HSP70 inhibits aminoglycoside-induced cochlear hair cell death and hearing loss. These data indicate that HSP70 induction is a critical stress response in the inner ear that can protect hair cells exposed to major stressors.

We utilized whole-organ cultures of utricles from adult mice to examine the mechanisms underlying the protective effect of HSP70. In response to heat shock, HSP70 was induced in glia-like supporting cells with little (or no) induction in hair cells. We developed a method of infecting supporting cells using adenovirus, and we utilized this technique to drive expression of HSP70 in supporting cells alone. Ad-HSP70 inhibited aminoglycoside-induced hair cell death, indicating that supporting cells are the primary mediators of the protective effect of HSP70. We hypothesized that HSP70 is secreted by supporting cells and internalized by hair cells. To test this, we performed a series of co-culture experiments using Transwell culture inserts. Co-culture of heat-shocked utricles with non-heat shocked utricles inhibited hair cell death in all utricles. Experiments using utricles from HSP70 knockout mice showed that the protective effect of co-culturing requires HSP70. In addition, depletion of HSP70 from the media using a function-blocking antibody abolished the protective effect in both heat-shocked and non-heat shocked utricles. These data support our prior data on HSP70 as a mediator of hair cell survival and suggest that HSP70 is secreted by supporting cells.

We propose that supporting cells can promote the survival of damaged hair cells. Taken together with recent data indicating that supporting cells may mediate hair cell death (Lahne and Gale 2008 J Neurosci 28, 4918; Bird et al., 2010 J Neurosci 30, 12545), our data suggest a major role for supporting cells in determining whether a hair cell under stress ultimately survives or dies.

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Damage-induced responses in cochlear and vestibular supporting cells: a Jekyll and Hyde story in which ERK1/2 is necessary and sufficient?
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Aside from certain cell types in the cochlea, the roles of inner ear supporting cells are not well defined. We know that during normal inner ear function supporting cells provide important structural support, but are also thought to provide critical trophic and homeostatic support for hair cells in both cochlear and vestibular systems. Work from a number of laboratories has shown that, under pathophysiological conditions, supporting cells are involved in repair of the epithelium and, in those systems that recover, supporting cells are the source of any replacement hair cells, whether it be by phenotypic conversion or a proliferative response. We have been studying the response of supporting cells during inner ear damage.

The extracellular regulated kinases 1 and 2 (ERK1/2) are activated in supporting cells surrounding an acute damage (laser/mechanical) site and also surrounding hair cells damaged by ototoxic aminoglycosides. Inhibition of MEK, the upstream kinase of ERK1/2, prevents ERK1/2 activation and affords some protection of inner hair cells in early postnatal cochlear cultures (Lahne & Gale 2008). In the chick utricule we have provided definitive evidence that nearest neighbour supporting cells are triggered to act as professional phagocytes. Moreover, the data suggested that supporting cells act not just as scavengers, but as potential mediators of hair cell death.

We now show that, in both neonatal and adult mouse utricules, ERK1/2 activation is also localized to the nearest neighbouring supporting cells during laser-targeted and aminoglycoside-induced hair cell damage. MEK inhibition (using U0126) in adult utricules resulted in a small but significant protection of hair cells after 24 hours of aminoglycoside treatment. The MEK inhibitor is not an ideal approach because long term application (24 hours) may well have negative effects on the hair cells themselves. Therefore, in order to determine the function of ERK1/2 activation in supporting cells we used an adenoviral infection protocol (Brandon, Voelkel-Johnson et al. 2012) to specifically infect supporting cells with a constitutively active (CA-) MEK1 construct. Using this to maximally activate endogenous ERK1/2 in supporting cells resulted in significant changes in cell behavior and a reduction in the number of hair cells compared to Ad-RFP infected controls. Moreover, live imaging revealed supporting cells expressing CA-MEK1 could be caught in the act of phagocytosing apparently healthy hair cells, indicating that ERK activation is sufficient to switch those cells from Dr Jekyll to Mr Hydes.

The enigmatic function of inner ear macrophages: Insights from studies of birds and mammals
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Macrophages are the first responders of the immune system and play essential roles in tissue maintenance and injury response. The sensory organs of the inner ear all contain resident populations of macrophages, which become activated and increase in number after hair cell injury. The avian cochlea contains numerous resident macrophages, which are mainly confined to the connective tissues and to the hyaline/cuboidal cell region. After hair cell injury, some macrophages migrate toward the damaged region, but they remain below the basilar membrane and do not enter the sensory epithelium. Experimental depletion of resident macrophages from the chick cochlea does not inhibit the removal of hair cell debris after ototoxic injury, and does not affect hair cell regeneration. However, macrophage depletion reduces the proliferation of cells immediately below the basilar membrane (the so-called mesothelial or tympanic border cells), suggesting that macrophages may serve a role in the maintenance of the basilar membrane.

In the mouse, the normal cochlea contains a small population of resident macrophages. Recruited monocytes from the vascular endothelium constitute the large majority of mononuclear phagocytes that populate the cochlea after sensory cell injury. Studies of Pou4f3-DTR mice suggest that the death of hair cells alone provides a sufficient stimulus to induce monocyte migration into the ear. As in the avian cochlea, the essential function of murine cochlear macrophages remains unknown. Macrophages and monocytes are abundant in non-sensory structures of the cochlea, and inflammatory cells are rarely observed interacting with hair cells or the sensory epithelium in animal studies. However, time-lapse imaging of mouse cochlear cultures have revealed that cochlear macrophages are intimately associated with spiral ganglion dendrites and can recognize and phagocytose apoptotic hair cells. Hair cell injury in vivo leads to increased macrophage
numbers within the osseus spiral lamina adjacent to the dendrites of afferent fibers as well as within the spiral ligament and limbus. The possible interaction between macrophages and afferent neurons is currently under investigation.

The vestibular maculae of chicks and mice also contain resident macrophages, which populate the stromal tissue below the sensory epithelium. Increased numbers of macrophages are recruited into the stroma of the chick utricle after aminoglycoside ototoxicity. Such cells often extend pseudopodia into the injured sensory epithelium, but they do not appear to play a key role in the removal of hair cell debris. As such, the function of vestibular macrophages remains a mystery.

### Using binaural detection to measure how precision of coding of interaural delay varies with both magnitude of interaural delay and center frequency

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**Background**

Explanations of binaural processing postulate that “internal delays” compensate for external interaural temporal delays (ITDs). Additionally, sound-field and earphone-based ITD-discrimination studies have consistently demonstrated that sensitivity to changes in ITD declines with increases in magnitude of reference ITD. Recently, we discovered a way to measure, quite directly, precision of ITD-coding as a joint function of magnitude of ITD and center frequency.

**Methods**

Binaural detection thresholds were measured in three conditions. The first consisted of transforming the classic NoSπ stimulus into an “(NoSπ):τ” stimulus by imposing an ITD on the entire signal-plus-masker waveform. With that stimulus, perfect internal compensation of external ITDs would yield thresholds both independent of ITD and equal to those obtained under NoSπ. Such compensation would transform (NoSπ):τ conditions back into NoSπ. Alternatively, if precision of ITD-coding declines with ITD, that type of transformation could not occur and thresholds would increase with ITD. The second condition capitalizes on the “double-delay” condition of van der Heijden and Trahiotis [(1999). J. Acoust. Soc. Am. 105, 388-399]. It is a variant of the (NoSπ):τ condition in which the masker is the sum of two independent noises, one interaurally delayed toward the left ear, one interaurally delayed equally toward the right ear. We call this (No)±:τ(Sπ):τ. For any τ, a double-delayed noise has the same interaural correlation as its “single-delayed” counterpart. There is, however, no internal delay that can be employed by the listener to transform (No)±:τ(Sπ):τ back to NoSπ and no value of internal delay expected to yield thresholds lower than could be achieved by imposing no internal delay. Thus, the difference, at any value of τ, between (No)±:τ(Sπ):τ and (NoSπ):τ thresholds reveals the detection advantage achieved by imposing a compensating interaural delay. The third condition employed is called (NoSo):τ and is a “monaural” control condition in which adding the signal to the masker produces no interaural cues and, thus, no binaural advantage.

**Results**

Data revealed that thresholds obtained using (NoSπ):τ stimuli centered at 500 Hz or 4 kHz increased with ITD and did so more rapidly at 4 kHz than at 500 Hz. The ordering of thresholds from low to high across τ was (NoSπ):τ, (No)±:τ(Sπ):τ, (NoSo):τ.

**Conclusion**

The data suggest strongly that precision of ITD-coding does, indeed, decline with ITDs up to, at least, 3 ms and does so more rapidly at 4 kHz than at 500 Hz.
head orientation (PHO), or auditory spatial mapping (ASM) changing as a function of roll angle. The aim of the current study was to distinguish between these two hypotheses.

Methods
Four participants were kept stationary at various roll angles of up to 120 degrees, where they lateralized noise bursts (150 ms, 0.3 – 12 kHz) presented simultaneously from two speakers. The speakers rolled with the participants and were near the interaural plane at azimuths of ± 37 degrees. A range of speaker level differences were employed to obtain psychometric functions and derive the speaker level difference corresponding to the AMP. No feedback was given. Simple equations for the PHO and ASM hypotheses were used to qualitatively model the results.

Results
The modelling data show that PHO and ASM vary linearly or sinusoidally, respectively, with roll angle. They also confirm that PHO, but not ASM, changes with the elevation of the sound source; for sounds along the interaural plane (zero elevation), PHO will not vary with roll angle. These modelling results suggest that experimental data from previous studies could not distinguish between the two hypotheses because head roll was limited to 90 degrees or less, and free-field sound sources were in the coronal plane. The current experimental data are consistent with the ASM-change hypothesis because AMP varies sinusoidally with roll angle, and the sounds were presented near the interaural plane. The experiment is currently being repeated with headphones, with sounds being perceived inside the head along the interaural plane. Preliminary results confirm that the direction of the audiogravic illusion is the same, further supporting the ASM hypothesis.

Conclusion
For free-field sounds, the audiogravic illusion can be explained by systematic changes in auditory spatial mapping (ASM). This would suggest that multisensory cues, predominantly vestibular otolith signals, directly influence the interpretation of binaural acoustic cues.

High frequency audibility and spatial release from informational masking
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Background
Previous work has shown that older and/or hearing-impaired listeners obtain less benefit of spatial separation in a multitalker informational masking experiment than do younger listeners with normal hearing. The goal of this study was to examine the role of high frequency speech information, which is often rendered inaudible by age-related hearing loss, in supporting spatial release.

Methods
A 20-dB range of target-to-masker ratios was examined in a brief test session, using the Coordinate Response Measure corpus. Two male masking talkers were presented across a range of levels and the task of the listener was to identify the color and number keywords spoken by a target male talker. Stimuli were adjusted for each listener to approximate the level above detection threshold that would be present for an ideal listener with no hearing loss (0 dB HL at all audiometric frequencies). Performance in this “broadband” condition was compared with a “lowpass” condition, in which the levels were adjusted to match the levels that would be present for an idealized hearing-impaired listener with a sloping high-frequency hearing loss (0 dB HL for .25, .5, and 1 kHz; -20 dB HL for 2 kHz; -40 dB HL for 4 and 8 kHz). Both the broadband and lowpass conditions were run at an RMS (before lowpass filtering) of 10 dB or 30 dB SL. SL was calculated relative to each listener’s speech detection threshold as estimated from that listener’s audiogram. A group of more than twenty listeners varying in age and hearing ability participated in the testing.

Results
Raising the target level from 10 to 30 dB SL generally improved performance, but, surprisingly, many of the hearing-impaired participants performed better in the lowpass condition than in the broadband condition. Many of the normal hearing listeners, on the other hand, performed better in the broadband condition, as would have been expected due to the increased information present. This was true at both sensation levels.

Conclusion
These data are not consistent with the hypothesis that older and/or hearing impaired listeners obtain less spatial benefit due to reduced audibility in the high frequencies. On the contrary, the data suggest that for some listeners with impaired
hearing there may be a disadvantage to high frequency energy, perhaps due to binaural interference or other types of distraction associated with the presence of energy in regions of impaired auditory function.

Modeling Sound Localization for Normal and Binaural CI-Listeners
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Background
Although modern cochlear implants (CI) provide good speech intelligibility, their performance in a “cocktail party” situation with multiple simultaneous sound sources is not satisfactory. It is well known that sound localization provides important cues to separate sources which enhance speech intelligibility in noisy environments. Since CIs were initially designed for monaural implantation only, there is room to optimize coding of binaural information.

Methods
Within a simulated listening setup the binaural Lindemann model can provide a metric to predict source localization abilities. After reviewing the model’s basic setup, we extended the model to work with action potentials. These were generated by physiologically inspired models which replicated auditory nerve responses for both intact ears and ears with CIs and different coding strategies.

The simulated listening setup consisted of a loudspeaker, which was circling around the listener’s head. The distance between the ears was 15 cm. An emitted wavefront reached both ears at different times and invoked interaural time differences (ITD). Furthermore, the simulation assumed linear air-attenuation and also generated interaural level differences (ILD).

The binaural Lindemann model extends the correlation delay line of the Jeffress model by inhibitory elements, thus adding ILD-sensitivity. A positive Lindemann correlation time delay indicates a sound source right hand side. A sound source with 1 Hz circling frequency will lead to a deviation of max. 0.441 ms.

Results
For a normal hearing model the Lindemann correlation easily tracked the sound source (supplement). When we tested MED-EL’s FS4 strategy, which conveys the temporal fine-structure in its four lowest frequency channels, we could also observe that the tracking of a moving source is possible, albeit with reduced precision (0.2 ms). When we analyzed the electrically induced spike trains, we found a much weaker Lindemann correlation due to the effects of channel crosstalk.

Conclusion
To improve sound localization of CI-listeners, coding strategies must convey ILDs and especially ITDs. We extended the Lindemann model to process input from spiking auditory models and used it for quantitative evaluations of localization cues of auditory models (normal hearing) and CI coding strategies. Whereas the CIS strategy did not provide useful fine-structure ITD cues, today’s latest coding strategies, the FS4 by MED-EL, was already able to convey viable fine-structure cues for sound source localization.

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Direction selectivity mediated by response adaptation in the owl's inferior colliculus

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Background

Direction-selectivity has been shown in the auditory system for frequency modulations and motion in space. The underlying mechanisms have been attributed to nonuniform topography in sensory representations and lateral connections. We investigated this issue in space-specific neurons of the owl's external nucleus of the inferior colliculus (ICx).

Methods

We recorded single units in the ICx of anesthetized barn owls. Moving sounds were simulated in a hemispherical free-field speaker array with 10° resolution in space. The owl was turned so that the RF of each cell was centered at 0° azimuth. We synthesized auditory drifting gratings, where the initial and final array configuration was constant at the beginning and end, irrespective of motion direction. The simulated moving sound started at one end of the array and moved around the owl's head at different velocities, in randomized directions. Spatial receptive fields (RFs) were measured in free field with randomized stimulation.

Results

Spatial RFs, as previously described, showed a main peak flanked by side peaks at 50-60° to each side. However, the sizes of the side peaks were often asymmetrical. The asymmetry can be caused by the convergence of ITD channels or by HRTF effects that attenuate the sound at certain locations. Surprisingly, most cells showed direction selectivity using the method described above. Direction selectivity calculated from RFs during simulated moving sounds strongly correlated with the size of the side peaks. Neurons preferred sounds coming from the side where the side peak was smaller. We found that the size of side peaks was correlated with their suppressive effects on the response to the center. Response to the side peaks also depended on the direction of motion. In addition, direction selectivity varied with velocity. These data are consistent with rapid excitation-induced response adaptation as the underlying mechanism.

Conclusion

Spatial receptive fields of direction-selective ICx cells can be strongly modulated by the direction of sound motion. Response adaptation acting on asymmetric spatial tuning can explain this selectivity.
Psychophysical Analogies to Auditory Source Width in Hearing-Impaired Adults
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Background
In a previous study, we demonstrated that older hearing-impaired (HI) listeners produced visual sketches of headphone-presented noises that were insensitive to changes in coherence. The current study further explores this insensitivity by presenting stimuli in the free field to HI and normal-hearing (NH) participants, and examines potential links in their perception of auditory source width to (a) binaural temporal fine-structure (TFS) resolution and (b) precision in absolute sound localization.

Methods
The stimuli for the sketching task were low-pass, high-pass and speech-spectrum noises. On each trial, a noise was presented from a center (0°) loudspeaker in a sound dampened room with two independent flanking noises that are simultaneously presented from loudspeakers at ±45° to act as arbitrary reflections. The flanking noises were attenuated 0-20 dB from the center signal level to generate partially coherent noises. HI and NH participants sketched the width of each stimulus presented repeatedly in random order. For the localization precision task, participants located 500-ms speech-spectrum filtered click trains presented from -30 to +30° in quiet. For the binaural TFS task, participants discriminated interaurally phase-shifted tones from diotic tones presented over headphones.

Results
The binaural TFS and localization precision data were in accord with previous studies showing the deleterious effects of age and impairment on these two measures, except that a portion of older participants could not perform the TFS task at all. When participants are parsed by their ability to do the TFS task, the sketching data showed a decreasing sensitivity to width with age and impairment that was not apparent in the headphone-presentation study: the worse one’s binaural TFS threshold – which was correlated with age – the less the perceived width changed with interaural coherence.

Conclusion
These results suggest that perceiving changes in the width of sound sources is related to the ability to hear changes in low-frequency temporal information. For those HI participants with binaural TFS data, the perception of sound sources was not wider, despite having decreased localization precision. This finding indicates that senescent changes to the auditory system do not necessarily lead to perceptions of broader, more diffuse sound images based on interaural coherence. Thus, the use of interaural coherence to predict sound-source perception, as often assumed in architectural acoustics, may not hold for older hearing-impaired individuals.
On the Integration of Binaural Spatial Auditory Cues and Self-motion (Idiothetic) Cues
W. Owen Brimijoin1, Michael A. Akeroyd1
1MRC Institute of Hearing Research

Background
Our acoustic world is in constant motion because the ears are attached to the head and the head is never perfectly still; we tend, however, not to notice this movement. Here we summarize experiments examining phenomena that appear to be related in that they each require an ongoing and highly accurate comparison of binaural spatial cues with ‘idiothetic’ cues (information on self motion derived from e.g. vestibular, proprioceptive, motor, or visual input). We will argue that these and related spatial phenomena are evidence of an auditory spatial processing mechanism that compensates for self motion.

Methods
We designed a system to move real and virtual sound sources as a function of head movements using a combination of a Vicon motion-tracking system (or Nintendo wii remotes) and real time digital signal processing. In the free field, we moved physical sound sources in a ring of loudspeakers, allowing us to create noises with illusory positions. In the free field and over headphones we moved sound sources in synchrony with head movements to examine the role of self motion in the externalization and internalization of sound sources. Finally, we used virtual sound sources that either moved with head movements or moved independently to examine whether there exists a fundamental difference between the processing of self motion and ‘real’ motion.

Results
We found that (1) signals that move predictably as a function of head movements can differ in perceived location from those that either remain static or move along arbitrary paths. (2) For noises, these illusory positions can affect their masking strength in a spatial release from masking paradigm. (3) Signals with fixed positions relative to the world are more likely to be perceived as being out in the world, whereas (4) those that move with the head tend to be internalized. Finally, (5) we found that self-generated auditory motion may be to some extent compensated for during moving spatial auditory tasks.

Conclusion
The results from each experiment suggest that auditory spatial perception is continually informed by self motion; that is, listeners constantly compare their own movement with the movement of the auditory world. All are consistent with the hypothesis that there exists a low-level integrative mechanism, not unlike the vestibulo-ocular reflex, whose function is to use idiothetic cues to adjust and stabilize auditory spatial representation in order to compensate for self motion.
Evaluation of a missing source identification paradigm for the assessment of spatial hearing ability in hearing aid users

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Background

When a sound source in a complicated auditory environment suddenly turns off, listeners with normal hearing often have the subjective impression that they can tell what the sound was and where it was coming from even in cases where they clearly were not paying any attention to the sound before it was removed from the scene. However, little is known about the extent to which hearing impaired listeners may be able to maintain awareness of multiple sounds in their environment, or the impact that hearing aids might have on this capability. In this study, we conducted a preliminary evaluation of a spatial awareness paradigm that required listeners to detect the addition or deletion of a realistic environmental sound from the auditory scene. The study was conducted with hearing-impaired bilateral hearing aid users under three listening conditions: bilaterally aided, monaurally aided, and unaided.

Methods

Isolated environmental sounds were presented from an array of 30 loudspeakers located in a cylindrical pattern spanning +/- 135 degrees in azimuth and +/- 30 degrees in elevation. Listeners were first asked to listen to a 2-s environmental sound and determine the identity of the sound and its spatial location. They then participated in additional blocks that required them to identify a source that was either added or deleted from the scene after a listening interval where two, four, or six environmental sounds were presented for 6 seconds from speakers at different azimuth locations. Each block was administered twice for each of the three hearing aid conditions over two sessions.

Results

Preliminary results revealed that hearing impaired listeners (n=7) had the most difficulty localizing the sound source in the unilateral-aided condition. Localization accuracy in azimuth was comparable in the un-aided and bilaterally-aided conditions, but source identification accuracy and localization accuracy in elevation were slightly better in the unaided condition than in the bilaterally-aided condition.

Conclusion

Preliminary results show a localization benefit for bilateral amplification over unilateral amplification, but no significant benefit for bilateral amplification over unaided listening. In the future, it may be possible to developed more advanced hearing aid algorithms that use binaural processing to enhance auditory awareness in hearing impaired listeners. The result of this experiment suggest that the missing source identification paradigm may provide a useful tool for evaluating changes in spatial awareness in associated with these new binaural hearing aid processing systems.

The Sound of Danger: Rapid Transmission of Auditory Stimuli to the Amygdala

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In the process of exploring the neural pathways through which the brain comes to associate auditory stimuli with aversive events we found evidence for direct inputs to the amygdala from the auditory thalamus. Behavioral studies demonstrated that these pathways are utilized in auditory threat (fear) conditioning. Physiological and molecular studies identified cellular and molecular mechanisms that underlie the learning. It is likely that thalamic inputs to the amygdala are allow for the creation of rapid but imprecise representations which are then refined via cortical inputs.

Morphological, neurochemical and connectional organization of the non-lemniscal thalamocortical pathways

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The function of the medial geniculate body (MGB) in normal hearing still remains largely enigmatic, in part due to the relatively unexplored properties of the non-lemniscal MGB nuclei. Indeed, the canonical view of the thalamus as a simple relay for transmitting ascending information to the cortex belies a role in higher-order forebrain processes. However, recent anatomical and physiological findings now suggest widespread roles for the non-primary auditory thalamic nuclei beyond that of a simple relay. Neurons in the non-lemniscal nuclei have distinct dendritic and somatic morphologies and
unique expression of calcium-binding proteins. Furthermore, the non-lemniscal nuclei send and receive feedforward and feedback projections among a wide constellation of midbrain, cortical, and limbic-related sites. Combined, these neuroanatomical features support roles for the non-lemniscal nuclei as conduits for auditory information flow to higher auditory cortical areas, mediators for transitioning among arousal states, and synchronizers of activity across expansive cortical territories.

**Ascending projections to the non-lemniscal auditory thalamus**

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It is generally assumed that the auditory thalamus receives ascending information predominantly from the inferior colliculus (IC) via three main parallel pathways: the central nucleus, dorsal cortex and external cortex of the IC (CNIC, DCIC and ECIC) innervate the ventral, dorsal and medial divisions of the medial geniculate body (MGBv, MGBd and MGBm), respectively. However, the literature provides conflicting evidence regarding the precise place of origin and termination of these pathways.

Using anterograde and retrograde tracers, we have investigated systematically the ascending projections to the auditory thalamus of the rat, with emphasis on the non-lemniscal nuclei. While our results confirm the dense CNIC-to-MGBv projection, they do not support the prevailing model representing the organization of projections from the DCIC and the ECIC. Specifically, the deep layers of the DCIC innervate the MGBv, much like the CNIC, but not the MGBd; the latter appears to be innervated only by neurons in the superficial layer 1 of the DCIC.

Perhaps our most unexpected result is that the MGBm and the suprageniculate nucleus do not receive significant projections from any IC subdivision. The main thalamic target of the ECIC is the posterior limitans nucleus (PLi), which extends for considerable rostrocaudal and dorsoventral distances as a narrow sheet along the lateral border of the anterior pretectal nucleus. Other significant targets of ECIC projections include the posterior intralaminar nucleus (PIN), the most dorsomedial region of the MGBd, the marginal zone, and the subparafascicular nucleus. These projections arise preferentially from different ECIC layers.

Our results also indicate that the auditory thalamus receives direct projections from an unexpectedly large number of neurons in subcollicular centers, including the contralateral dorsal cochlear nucleus and lateral superior olive, the ipsilateral medial superior olive, superior paraolivary nucleus, and ventral nucleus of the lateral lemniscus, and the dorsal nucleus of the lateral lemniscus and the nucleus sagulum of both sides. Some of these projections target preferentially the MGBm.

In summary, the ascending projections to the auditory thalamus follow several pathways that do not necessarily go through the IC. The MGBm participates in a fast track from subcollicular centers that bypasses the IC and transmits converging information to the cerebral cortex. On the other hand, the auditory information from the non-lemniscal regions of the IC is not conveyed to the MGBm, but to other thalamic nuclei, chiefly the PLi and the PIN, presumably to subserve acousticomotor and emotional functions.

**Response properties of non-lemniscal MGB neurons to auditory stimuli in vivo**

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The principal nucleus of the auditory thalamus, the medial geniculate body (MGB), can be divided into ventral, medial, and dorsal subdivisions. Traditionally, the ventral division is the only subdivision considered to be part of the lemniscal or primary auditory pathway, while the medial and dorsal subdivisions are assigned to the non-lemniscal or secondary pathway. Based on physiological properties of neurons recorded from histologically identified subdivisions within the MGB of anaesthetised rodents, I suggest medial MGB neurons do not fit simply into the conventional functional dichotomy of high-fidelity lemniscal, and context-dependent non-lemniscal pathways. For example, while a high proportion of medial MGB neurons display change sensitivity (a property typical of non-lemniscal areas), neurons in the medial MGB also possess response fidelity (short latencies, high response probability, and an ability to follow sounds with high rates of temporal modulation) that frequently exceeds that of neurons present in the (lemniscal) ventral MGB. This enhanced reliability may be inherited from direct inputs from the auditory brainstem – a pathway unique to the medial MGB. Taken together, these results argue in favour of a distinct role for medial MGB in central auditory processing.
Computational, physiological and anatomical findings that distinguish MGBm processing
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BACKGROUND: The medial region of the medial geniculate body (MGBm), broadly defined to include the medial division of MGB, the suprageniculate and the posterior intralaminar nuclei, surrounds the ventral (MGBv) and dorsal (MGBd) divisions of the MGB on their ventral and medial aspects. MGBm is the most heterogeneous region of the MGB, in terms of connectivity, anatomy and physiology. Unlike MGBv and MGBd, MGBm projections mainly avoid middle cortical layers or they project to subcortical regions, where they have been shown to be critical for the formation of auditory-based associative memories. Here some salient differences in anatomy and physiology will be reviewed, along with new computational and anatomical data demonstrating how those differences are likely to influence auditory processing.

METHODS: Single compartment models of MGBm neurons are created using NEURON software. These models incorporated physiological data obtained from rat brain slice studies, including membrane properties, excitatory synaptic properties, and inhibitory synaptic properties. In addition, the distributions of ascending axon terminals from the IC are estimated using vesicular glutamate transporter 2 (vGluT2) immunohistochemistry.

RESULTS: Physiological results have shown that MGBm neurons often lack a low-threshold calcium current, a hallmark of standard thalamocortical neurons. In addition, many of these neurons have an apamin-sensitive calcium-activated potassium channel that causes MGBm neurons to adapt to sustained current injections. These properties are incorporated to compare model responses to collicular inputs in MGBm versus MGBv neurons.

CONCLUSIONS: Many MGBm neurons should not be grouped with MGBv and MGBd neurons when considering their physiology or their participation in thalamocortical sensory processing.

Dynamic sound representations in the lemniscal and extra-lemniscal divisions of the medial geniculate body
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The medial geniculate body (MGB) of the thalamus occupies a critical node between corticopetal, corticofugal and limbic processing networks. The corticothalamic (CT) projection from layers 5 and 6 (L5 and L6, respectively) represents the largest component of the auditory corticofugal network, and one of the largest projection systems in the brain. CT feedback has been implicated in a diverse range of cognitive functions ranging from memory consolidation to dynamic sensory filtering, yet a detailed understanding of its functions and mechanisms has remained elusive due to the technical difficulties associated with selectively manipulating CT neurons without disrupting thalamocortical (TC) or intracortical processing. We found that CT axon terminals were restricted to a circumscribed zone within the dorsal subdivision of the MGB (MGBd) shortly after birth but had invaded surrounding regions in the ventral and medial subdivisions (MGBv and MGBm, respectively) by the time of hearing onset. Using in utero gene electroporation, we over-expressed EphA7, a receptor tyrosine kinase, in the auditory cortex, and were able to forestall the developmental elaboration of CT axonal projections into the MGBd and MGBm without disrupting TC projections. Electrophysiological recordings from the MGBv of adult mice with chronically reduced CT feedback revealed lower baseline spike rates compared to controls yet surprisingly subtle effects on sound-evoked response properties. Our most recent experiments explore the effects of acutely reducing CT feedback. Using a chemical-genetic approach, we were able to selectively and reversibly inactivate CT neurons in a single cortical layer while recording from ensembles of neurons in lemniscal and extra-lemniscal divisions of the MGB. We found that activating the inhibitory DREADD, hM4d, eliminated tonal receptive fields and strongly reduced firing rates in genetically targeted cortical neurons within 15 minutes and lasted for several hours. In contrast to the modest effects of chronic CT reduction, our ongoing experiments suggest that acute silencing of L6 cortical neurons via hM4D can strongly modulate sound-evoked responses in the MGB of awake, passively listening mice.

Interactions between auditory and cognitive aging
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Background: Auditory-cognitive interactions are important because the mutual influences of bottom-up and top-down processing shift depending on the listener, signal, task and context, and also because there can be brain re-organization
to compensate for sensory and/or cognitive impairments. In cognitive aging, knowledge and use of context is preserved, but processing (working memory, attention/inhibition, speed of processing) declines. There are at least three sub-types of age-related hearing loss, including neural-type with little audiometric loss.

The trio of working memory, attentional control and speed of perceptual processing are inter-related. This trio operates differentially depending on bottom-up factors, including the quality of the speech signal and the nature of hearing loss. Methods: Auditory-cognitive interactions will be illustrated with an overview of experiments in younger and older adults where the speed and/or accuracy of speech understanding is altered by manipulating the signal quality, contextual cues that enhance attention (semantics, emotion, location, timing, talker voice), or task demands with varying cognitive processing loads. Context is manipulated in experiments in which 1. materials vary in the congruency of semantic context to facilitate or inhibit the speed of lexical decision; 2. the vocal emotion of the talker modulates word recognition accuracy in noise; 3. word recognition accuracy in a multi-talker scene varies depending on whether the location of the target is likely or unlikely; 4. vowel recognition in noise depends on the relationship between the timing of speech and expectancy based on musical stress patterns; 5. word recognition accuracy in noise develops with streaming based on talker voice. Cognitive load post word recognition is varied in terms of how the listener must manipulate the words and/or how many words are to be recalled.

Results: The illustrations demonstrate that there are both auditory effects on cognition and cognitive effects on audition. Age-related and individual differences suggest that there are short- and long-term, and inter- and intra-individual variations in auditory-cognitive interactions. Unifying themes across phenomena will be proposed.

Conclusions: Implications for research and practice will be suggested, including a new perspective on the possible short- and long-term benefits to both hearing and cognition of a rehabilitative approach that marries improving signal quality and training to increase active engagement in listening. This overview talk will set the stage for the following presentations in the session.

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Individualization of hearing aid signal processing based on cognitive measures

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Hearing instruments use complex processing anticipated to improve speech understanding. While many listeners benefit from such processing, others do not and these differences are potentiated by adverse listening conditions. Unwanted artefacts of hearing aid signal processing may contribute to adverse listening conditions (Mattys et al., 2012). Under such conditions cognitive processing resources are allocated to the recovery of degraded information at the auditory periphery, leaving fewer resources for understanding the message.

Working memory refers to the capacity to hold and manipulate a set of items in mind, such as speech fragments. A number of recent studies indicate that individual working memory capacity influences the individual benefit of hearing aid signal processing. Ng et al., (under revision) examined the effects of binary masking noise reduction (Wang et al., 2009) in hearing aids on memory processing of speech perceived in a competing speech background, using a sentence-final word identification and recall test. The reading span test was also administered. The reading span test assesses the simultaneous processing and memory aspects of working memory. Noise reduction lowered the negative effect of noise on memory performance. Participants with good working memory capacity obtained a further benefit. Lunner and Sundewall-Thorén (2007), among others, showed that the individual benefit of fast wide dynamic range compression was associated with individual working memory capacity. Persons with high working memory capacity benefited most from fast acting compression while the persons with low working memory capacity benefited most from slow acting compression. Arehart, Souza, Baca and Kates (in press) showed that elderly individuals with low working memory capacity were more disadvantaged by frequency compression processing during speech recognition in noise than elderly individuals with high working memory capacity. It is possible that the negative effects of fast acting wide dynamic compression and frequency compression for persons with low working memory are due to unwanted artefacts of signal processing.

In conclusion, the benefits of certain given implementations of complex processing in hearing aids have to be weighed against the extra cognitive load that such processing seems to engender, possibly as a result of unwanted artefacts. Although complex processing may aid speech understanding in noise for individuals with hearing impairment and especially those with good working memory capacity, it seems that for some individuals with lower working memory capacity, the drawbacks may outweigh the benefits. This should be taken into account when hearing aids are fitted.
Processing load during speech perception in noise – insights from pupillometry
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Title: Processing load during speech perception in noise – insights from pupillometry
Authors: Adriana A. Zekveld, Thomas Koelewijn, Sophia E. Kramer.

Background
Pupillometry provides an objective method for measuring cognitive load (listening effort) during speech perception in adverse listening conditions. We assessed the influence of individual factors like age, hearing loss, and cognitive ability and external factors like stimulus type, stimulus modality, intelligibility level, and masker type on the pupil response.

Methods
Combining and comparing the results of 4 studies allowed us to assess the influence of speech intelligibility level and masker type on cognitive processing load during listening and how this relates to cognitive ability. Five groups of young ($n_{tot} = 123$), two groups of middle-aged normal-hearing subjects ($n_{tot} = 62$) and one group of middle-aged hearing-impaired subjects ($n_{tot} = 38$) participated. Pupil dilation was recorded during speech perception in noise and during the text reception threshold (TRT) test. Sentences and words were presented over a wide range of sentence intelligibility levels (1% - 99% correct), masked by different masker types (steady-state, fluctuating, single-talker) and in quiet.

Results
The results show consistently larger pupil responses in less intelligible or more complex listening conditions. The pupil response increases with decreasing speech intelligibility level. However, when speech perception is very difficult and subjects start to give up trying to perceive the speech, the pupil response decreases relative to less difficult conditions. This inverse-U shaped function of the pupil response across intelligibility levels supports the validity of the pupil response as measure of cognitive processing load. Middle-aged and hearing-impaired adults show a smaller decrease in pupil response with increasing intelligibility. Independent of intelligibility level, masker type affects cognitive processing load. Processing load is highest for speech masked by an interfering speaker. The TRT test is associated with cognitive load during listening, especially in relatively difficult listening conditions. The data suggest that an individual’s cognitive abilities modulate the relation between speech intelligibility and processing load. The pupil response furthermore reflects processing load differences caused by different stimulus types (word versus sentence perception), task characteristics (detection versus identification) and stimulus modality.

Conclusions
The pupil response and speech perception performance data reflect different aspects of processing load during listening. The impact of several factors and individual differences that affect the complexity of the listening task and cognitive processing load are quantifiable by assessment of the pupil response during listening.
Listening effort and cochlear implants
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Background
Sound transmission through cochlear implants (CIs), although restoring hearing for deaf individuals, is far from perfect. Typically, speech transmitted through the device maintains coarse spectral and temporal information but lacks temporal fine structure. Interpreting this degraded auditory signal can be effortful, and CI users do anecdotally report fatigue. Therefore, in addition to increasing speech intelligibility, a purpose of CIs could be to reduce listening effort. However, to be able to do so, a reliable measure of effort is needed. In this study, we proposed to use a dual-task paradigm to quantify listening effort with CIs. We hypothesized that listening effort may change differently than intelligibility for different CI programs, which may not be reliably captured with traditional speech intelligibility tests.

Methods
The dual-task paradigm is based on the idea that cognitive resources are limited and effort of the first task can be quantified by changes in performance on the secondary task. In the present study, the primary task measured the intelligibility of sentences processed with a noiseband vocoder simulation of CIs. The secondary task, rhyme judgment, measured the response times (RTs). In the first experiment, using this paradigm, intelligibility and RTs were measured for a varying number of spectral channels of CI simulations with normal-hearing native Dutch speakers. In the second experiment, with a similar population, intelligibility and RTs were measured for a number of simulated CI configurations, such as a single CI and bimodal hearing of a CI combined with a hearing aid.

Results
Experiment 1 showed that speech perception, measured in percent correct, plateaued at 6 channels while listening effort, measured in RTs, further decreased up to 8 channels. Preliminary results from Experiment 2 did not show a significant reduction in effort with simulated bimodal hearing over a simulated single CI.

Conclusions
Results from the first experiment have shown that listening effort may be further optimized even after seemingly perfect speech perception is achieved, confirming our hypothesis. Preliminary results from the second experiment have not yet indicated an improvement in effort with bimodal hearing simulations, however this may have also been caused by not selecting the right parameters in the dual-task paradigm. Further research is conducted to pursue listening effort with new paradigms using cochlear implant simulations, as well as actual implants, and different implant configurations.
Measuring Listening Effort with Reaction Time to Digits in Noise
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Background: Audiological rehabilitation can be beneficial for speech communication even if it does not have a measurable effect on speech intelligibility. For example, single-channel noise reduction is often preferred by hearing-aid wearers, while it does not improve speech intelligibility. We hypothesized that this is caused by an improvement in listening effort. Unfortunately, there is no easy method for the measurement of listening effort. The purpose of the current study is to determine if the measurement of the reaction time to spoken digits in noise can be used to measure listening effort at signal to noise ratios (SNRs) that are high enough to render the speech completely intelligible. Additionally we want to determine if the reaction time to spoken digits can be improved by noise reduction processing.

Methods: We added various amounts of stationary noise to spoken digit triplets to obtain different SNRs (-10 dB, -5 dB, 0 dB, quiet). In addition to these "unprocessed" speech-in-noise stimuli we also included stimuli that were processed with two noise reduction schemes (an ideal binary mask and a Wiener algorithm with MCRA-noise estimator). All stimuli were used in a reaction time experiment with two different tasks. In the first task, participants had to quickly identify the last digit ('identification'). In the second task they had to quickly add the first and the last digit ('arithmetic'). Twelve normal-hearing participants participated in this study.

Results: At all SNRs the response time was increased significantly for lower (i.e. worse) SNRs for both the identification and the arithmetic tasks. The response time on the arithmetic task was more influenced by the noise than the response time on the identification task, but the arithmetic task led to a higher variance. Measurement of the effect of noise reduction on reaction time is ongoing.

Conclusions: The significant effect of SNR on reaction time implies that it is possible to measure listening effort with spoken digits at SNRs at which speech is highly intelligible. The optimal task may depend on the SNR that is of interest. A potential audiological application is the evaluation of noise reduction and data investigating this application are currently being gathered. If positive, this listening effort test would fill a gap in the evaluation of assistive hearing devices.

Seeing in the dark: MEMRI of the neuroretina
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Functional MRI (fMRI) is usually based on detectable hemodynamic changes during a stimulus (BOLD). This endogenous contrast mechanism has greatly facilitated fMRI's wide application; however, the information about neuronal activity generated by BOLD studies is indirect and limited to identifying the spatial location and spatial extent of activity. Here, I describe our studies using the neuroretina as a model system for evaluating another functional imaging approach, manganese-enhanced MRI (MEMRI). Importantly, MEMRI not only pinpoints functionally active regions but also allows for an analytical assessment of the degree of neuronal activity (based on calcium ion influx pathways like L-type voltage gated calcium channels) in that localized region-of-interest. The potential of applying MEMRI to translate studies between bench and bedside and back will also be discussed.

Detecting Laminar Specific Neural Plasticity in the Rodent Brain with MEMRI
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Detecting Laminar Specific Neural Plasticity in the Rodent Brain with MRI
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Functional MRI techniques have found widespread use to measure brain neural circuits that are used for a large number of behaviors. When circuit activity changes due to plasticity, it remains a challenge to identify sites of synaptic changes responsible for the circuit level changes measured. Of particular interest are the cases of long range cortical rearrangements that have been detected in the human brain after injury. Rodent models that mimic some of these cortical rearrangements have been developed and are being used to determine the synaptic basis for the plasticity detected. We have developed a model of adult cortical plasticity due to peripheral somatosensory nerve damage that is being used to develop MRI tools that can pinpoint sites of synaptic changes. Two weeks after peripheral denervation of one side of the forepaw, hindpaw, or whisker pathway there is a large up-regulation of cortical activity from the spared side and a large...
up-regulation of callosal inputs from the spared cortex to the cortical representation of the denervated area. A combination of functional MRI and laminar specific neural track tracing using manganese enhanced MRI predicted changes in thalamo-cortical inputs to layer IV that contribute to the up-regulation of cortical activity along the spared whisker barrel pathway. Slice electrophysiology confirmed that the thalamic inputs on layer IV stellate cells were strengthened by a post-synaptic mechanism. Sites of plasticity that explain the up-regulation of the callosal communication have also been studied with MRI. High temporal-spatial resolution fMRI demonstrates that up-regulation of the communication between the spared and denervated cortices likely occur through callosal inputs. These fMRI results were consistent with manganese enhanced MRI that predicts a strengthening of inputs into layer 2/3 and 5. Taken together these results demonstrate that MRI is positioned to begin to give laminar specific information about mechanisms of cortical plasticity.

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**Studying development and plasticity of auditory function with MEMRI**

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There are several challenges to studying the emergence of central auditory processing. The sound-driven responses of juvenile neurons are small in magnitude and adapt rapidly. Furthermore, these characteristics can be exacerbated by the anesthetics the permit stable recordings. As a consequence, studies of functional development are usually based on a relatively sparse sample of neurons, and the effect of developmental manipulations, such as hearing loss or abnormal stimulation, are usually assessed in adulthood when robust recordings can be more easily obtained. In this presentation, I will review the strengths and limitations of Manganese-enhanced magnetic resonance imaging (MEMRI) for the purpose of studying auditory processing in control and experimentally reared animals.

Mice were raised either under normal laboratory conditions, or with conductive hearing loss, or in a specific acoustic environment. Each animal was then injected with MnCl₂, and Mn²⁺ uptake was induced by exposure to either a broadband noise or a single pure tone while awake. Animals were imaged in an anesthetized state using a 7-Tesla horizontal magnet and 25mm head coil. A 3D gradient echo pulse sequence was used to acquire T1-weighted brain images, with an isotropic spatial resolution of 100 μm. Volumetric image data from each brain was registered to a 3D template, and the inferior colliculus (IC) was extracted using an mask segmented from the template. 3D images were analyzed statistically with a voxel-wise multiple-comparison method in Matlab (see Yu et al. 2005, 2007, 2008).

Three sets of developmental findings were obtained with MEMRI. First, the 3D map of sound frequency in IC displays significant refinement during the week after hearing onset. Second, developmental exposure to sound frequencies that are ordinarily represented in IC by non-overlapping volumes, causes a significant reorganization of the 3D map. Third, the effect of conductive hearing loss on IC activity depends on the age of hearing loss onset and duration of deprivation.

MEMRI offers certain advantages for studying functional development. Stimulus-specific activity patterns can be induced in awake animals, and the measurement obtained subsequently from all brain regions with a high spatial resolution. Furthermore, this activity pattern can be obtained, noninvasively, from the same animal at a later age. However, MEMRI activity patterns provide no information about neuron discharge patterns that encode specific sound cues. Therefore, MEMRI is best employed when whole brain mapping and assessment points are required to address a hypothesis.

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**Noise-induced Calcium-dependent Activity in the Central Auditory Pathway: A Manganese-Enhanced MRI Study**

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**Background**

Noise exposure at high intensities leads to a temporary shift of hearing thresholds (TTS) and is followed by a permanent threshold shift (PTS). Permanent threshold shift is not only associated with cochlear damage as the primary site-of-lesion, but also with subsequent structural and functional changes within the central auditory pathway. The aim of the present study was to monitor neuronal activity within central auditory structures in mice after noise exposure at different time intervals using manganese-enhanced magnetic resonance imaging (MEMRI).

**Methods**

Young adult mice were noise exposed for 3 hours with a broad-band white noise (5–20 kHz) at 115 dB SPL. Twenty-four hours before the MRI-scanning was performed using a 7-Tesla rodent scanner, a 0.4 mM/kg dose of MnCl₂ was injected.
Mice were investigated immediately after the noise exposure, or were kept in their cages for one or two weeks to be investigated at 7 or 14 days after noise exposure. Another group was noise-exposed for a second time at one week after the first exposure. These animals were investigated 7 days after the second exposure. One additional group was exposed to sound intensities of 90 dB SPL (moderate noise exposure) and investigated 7 days later. Unexposed animals were used as controls.

**Results**

Calcium-dependent activity patterns were modified in several structures of the central auditory system as the result of noise-induced hearing loss (NIHL). The MEMRI-data demonstrate that temporary threshold shift is correlated with an activity increase in hierarchically lower structures of the auditory pathway. This seems to be indicative of a direct noise impact at the first stage of central auditory processing. However, noise-dependent changes of higher auditory structures were found as well in the phase of PTS. Repeated noise exposure was found to induce an additional elevation of calcium-dependent activity in all investigated auditory structures — without a significant shift in auditory thresholds. Sustained manganese accumulation was present in the auditory brainstem after moderate acoustic stimulation as well without PTS induction.

**Conclusions**

The long-lasting enhancement of MEMRI signals suggests a noise-induced activity increase of various calcium-dependent processes of different origin. The present findings could be helpful to better understand the time-course of different symptoms in NIHL and the individual susceptibility to noise.

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**Manganese Enhanced MRI (MEMRI) to Assess Axonal Transport Deficits and Improvements in Mouse Models of Alzheimer’s Disease**

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**Background:** Axonopathy is observed in many neurodegenerative diseases. In Alzheimer’s disease (AD), axonal swellings and degeneration are prevalent and are thought to contribute to AD pathology. Current limitations in identifying the contribution of axonal damage to AD include the inability to detect when this damage occurs in relation to other identifiers of AD because of the invasiveness of existing methods. To overcome this, we used the MRI methodology Manganese Enhanced MRI (MEMRI) to assess *in vivo* axonal transport rates in the Tg2576 mouse model of AD and in Tg2576 mice crossed with superoxide dismutase 2 (SOD2) overexpressing animals.

**Methods:** Mice were imaged using a 9.4T, 21 cm bore horizontal scanner with a 35 mm volume resonator (Bruker, BioSpin). Mice were anesthetized with 5% isoflurane mixed with oxygen and maintained on 1.5 – 2 % isoflurane during imaging. The imaging parameters to acquire olfactory multi-spin/multi-echo MEMRI images were as follows: TR=500 ms; TE=10.2 ms; FOV=3.0 cm; slice thickness= 1 mm; matrix=128°—128; NEX=2; number of cycles=15; using Paravision 5.0 software (Bruker BioSpin, Billerica, MA). The core temperature was maintained at 37 °C during scanning. The region of interest (ROI) was identified and placed on an axial slice 1 mm in front of the posterior of the olfactory bulb (OB). The ROI was determined by measuring the length of the olfactory bulb, locating the midpoint of this line, then extending this midpoint out to the ONL.

**Results:** Our results demonstrate that axonal transport deficits are present in the Tg2576 mouse during amyloid beta accumulation and also before plaque formation. Furthermore, our results demonstrate that by reducing oxidative stress by overexpressing SOD2 in the Tg2576 mouse rescues axonal transport deficits.

**Conclusions:** MEMRI can be used to measure axonal transport deficits and improvements in mouse models. We have shown that axonal transport deficits are present in mouse models of AD before plaque formation occurs. We also show that this axonal transport deficit can be rescued by reducing oxidative stress through SOD-2 overexpression. The use of MEMRI to assess transport deficits and improvements should be applicable to additional mouse models of neurodegeneration.
New approaches for studying tinnitus: bridging the gap between basic science and current clinical concerns
Anthony Castracane
Wayne State University
Within the last decade, advancements in the study of tinnitus have been made by combining non-invasive neuroimaging modalities (anatomical and physiological) with non-invasive treatment paradigms. One such treatment paradigm involves the use of repetitive transcranial magnetic stimulation (rTMS) as a strategy for tinnitus abatement. Voxel-based morphometry (VBM) has been used as an anatomical outcome measure to evaluate plastic changes in the brain following treatment. Magnetic resonance spectroscopy (MRS) has also been used as a physiological outcome measure to evaluate changes in metabolites following rTMS treatment.

Recent experimental work has shown that manganese enhanced MRI (MEMRI) can identify tinnitus related neural activity in the nervous system in animals with behavioral evidence of tinnitus. To extend this exciting work, we now propose that manganese encapsulated nanoparticles may be the next advancement in tinnitus research when considered under the rubric of “nanotheranostics;” a new area of research that focuses on the combination of diagnostic detection with therapeutic drug delivery carriers. It is hypothesized that nanoparticles capable of housing manganese ions for MRI contrast enhancement, binding target-specific receptors, and releasing appropriate therapeutic agents in tinnitus positive brain areas, will improve treatment efficacy and ultimately lead to the best possible outcomes for individuals with tinnitus. Indeed, it is the functionality of nanoparticles that has inspired the use of this theranostic platform. In addition, because many current methods used for tinnitus abatement are either ineffective or produce inconsistent and sometimes disappointing results, new and innovative approaches are needed. Developing unique nanotechnology provides the platform and opportunity to attack the issue of tinnitus suppression head on. If successful, we can bridge the gap between basic science and the translational clinical potential to humans in order to make new in roads for novel drug treatments. In combination with other technology and imaging techniques, we are on the road to a cure for abating this often debilitating condition.

In vivo identification of hyperactive neural activity in rats with behavioral evidence of tinnitus using MEMRI
Avril Genene Holt, David Bissig, Najab Mirza, Gary Rajah, Bruce Berkowitz
Wayne State University School of Medicine
Currently, there are no drugs that are broadly effective in the treatment of tinnitus. Crucial for development and testing of pharmacological interventions is a biomarker that is a. reliably correlated with the perception of tinnitus b. apparent regardless of hearing status or method of tinnitus generation and c. persistent, present as long as tinnitus percepts remain. Many tinnitus studies have suggested that changes in spontaneous neuronal activity may be one such biomarker. A novel non-invasive method for assessment of neuronal activity is manganese enhanced MRI (MEMRI). In this study our group has used MEMRI to evaluate neuronal activity in auditory related nuclei. Two complementary methods were combined to evaluate established tinnitus models: inhibition of acoustic startle and manganese-enhanced MRI. In order to induce tinnitus, adult male rats were either administered salicylate once daily for three days or exposed to a single noise event (10 kHz, 118 dB one-third octave band tone, four hours). Salicylate-induced tinnitus (as determined with acoustic startle) resulted in wide spread manganese uptake that was significantly elevated when compared to uptake levels in both non-tinnitus rats and rats with acute noise-induced tinnitus sustained. Only in the inferior colliculus (IC) did both tinnitus models exhibit manganese uptake that was significantly elevated when compared to non-tinnitus animals. Therefore, abnormal neuronal activity levels in the IC appear to be important in tinnitus-mediated activity. Our results suggest that increased spontaneous neuronal activity in the IC is an indicator of tinnitus perception and that modulation of spontaneous neuronal activity may be used as a method for evaluation of proposed therapeutic intervention. These studies provide the foundation for future studies correlating the longevity of tinnitus with hearing loss and neuronal activity in specific brain regions resulting in tools for evaluating treatment efficacy across tinnitus paradigms.

Biofabricated manganese encapsulated nanoparticles: Novel theranostic applications to identify and treat tinnitus
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One challenge associated with targeted imaging and theranostic nanotechnology is the packaging and site-specific delivery of therapeutic/imaging agents to the tissue of interest. One route to a multifunctional nanoparticle platform is to
leverage existing biological systems capable of self-assembly and multivalent display. Bacteria and plant virus capsids are ideal candidates in this respect, where entrapment of dyes/drugs inside the capsid combined with external capsid modification offer a novel concept for nanoscale packaging and site-specific targeting. The icosahedral MS2 bacteriophage is a potential candidate for such a system, having an exterior diameter of ~28 nm with large pores, where small molecules to enter the capsid interior. MS2 also has several lysines available on the exterior for conjugation of targeting ligands. We have previously demonstrated RNA-based self-packaging of a heterocyclic photoactive agent (meso-tetrakis(para-N-trimethylanilinium)porphine, TMAP) into the MS2 capsid and used this system for targeted delivery of drugs to cancer cells. In the current work, we investigate the potential of the MS2 capsid for use in a similar fashion to encapsulate Mn-chelating agents applied to MRI imaging. Extended functionality by modifying the capsid exterior with targeting moieties will be discussed to allow targeted imaging/treatment for tinnitus research.

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Impact of short-term auditory training on subcortical processing of sounds
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Background: Auditory thalamus and brainstem structures play an important role in auditory learning. However, our understanding of the extent to which subcortical contribution is experience-dependent has been limited, particularly in healthy young adults. The purpose of our studies was to further our understanding of the neural impact of training on the subcortical processing of sound.
Method: In several studies, we investigated the impact of cognitive-based auditory training programs on perceptual ability and subcortical processing of speech via EEG and fMRI.
Results: Our findings show training-related perceptual improvement. In addition, neural recordings demonstrate continued malleability of subcortical encoding of speech in adulthood. Predictive relationships were also found between pre-training neural activity and amount of perceptual improvement, thus offering a useful metric to help identify individuals who would benefit most from auditory training.
Conclusions: Our studies offer empirical evidence that auditory training involving active engagement of cognitive processes can alter the neural encoding of sound by enhancing the processing of acoustic cues that are important in speech perception. Also, by demonstrating training-related improvements in normal hearing, young adults, we were able to better delineate the capacity for change as well as limitations of the central auditory system.
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Circuit organization and synaptic plasticity of neurons in the top-down auditory brainstem projecting pathway
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Top-down corticofugal pathways exert significant influence on auditory stimulus processing in the brainstem. The inferior colliculus (IC) is a site of synaptic convergence from other auditory brainstem nuclei; its role in transmitting this convergent information to the thalamus makes it an important target of top-down modulation. The function of top-down pathways targeting the IC have been studied using behavioral assays and by characterizing physiology at the level of the IC. However, little is known about how cortical networks at the origin of this top-down pathway are structured and activated.
To better understand this aspect of top-down auditory pathways we studied the microcircuit architecture and synaptic dynamics of IC projecting (cortico-collicular) neurons in the auditory cortex. Cortico-collicular neurons were labeled by injecting retrograde fluorescent tracers in the IC. We used laser-scanning photostimulation (LSPS) with glutamate uncaging to characterize the microcircuit architecture of monosynaptic inputs to IC-projecting neurons in layer 5B (L5B).
To assess the specialization of top-down circuits we injected fluorescent tracers in the contralateral auditory cortex and characterized synaptic input maps of L5B cortico-cortical neurons. We demonstrate that cortico-collicular and cortico-cortical networks form the basis of two functionally distinct auditory projection pathways. For example, cortico-collicular neurons receive stronger inputs from layer 2/3 than do cortico-cortical neurons. To understand the synaptic dynamics of this pathway we performed dual whole cell recordings on connected L2/3 and L5B neurons. We found that L2/3→L5B cortico-cortical synapses obeyed conventional short-term plasticity rules: stronger synapses depress with repetitive activation and weaker synapses facilitate. In contrast, we observed a different form of plasticity at L2/3 synapses projecting to cortico-collicular L5B neurons: weaker synapses depress more than stronger ones. Computational models of an auditory cortical column allowed us to test the functional implications of these differential forms of plasticity. Short-term synaptic plasticity in the cortico-cortical column effectively shuts down the L2/3→L5 pathway, and by consequence L5 cortico-cortical output.
Short-term synaptic plasticity maintains large-amplitude L2/3->L5 synapses thus keeping the cortico-collicular pathway open after short term sensory activation. Our findings establish that cortico-collicular and cortico-cortical networks form distinct parallel functional pathways in the cortex and we expect our findings will establish a new framework for the study of top-down networks in the auditory system.

**The disadvantaged brain: Auditory function and socio-economic standing**

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**Background:**
Studies of auditory deprivation in humans typically focus on the effects of deafness. However, deprivation can occur when hearing thresholds are intact. For instance, children from families of low socioeconomic standing (SES) hear fewer words per hour, receive less cognitive stimulation, and face greater noise exposure than their high SES counterparts. Despite the prevalence of poverty in the United States, little is known about how this early adversity affects the human brain. What is known, however, is that parental SES is a strong predictor of a person's life trajectory and that environmental factors, including auditory deprivation, can have lifelong repercussions. From this, we propose that auditory function and SES interact reciprocally.

**Methods:**
To characterize how SES affects auditory brain function, we compared two groups of normal-hearing adolescents that attended the same schools and were matched in age, sex, and ethnicity, but differed in their maternal education level (11.43 vs. 15.00 years), a correlate of socioeconomic status. We recorded auditory brainstem responses (ABRs) to the speech syllable "da" in addition to measuring neural noise in the absence of stimulation. From the speech-evoked ABR, we measured the fidelity of the response to the vowel (spectral power within 284-655 Hz) and computed within-session response consistency by comparing the first and second halves of the recording.

**Results:**
Adolescents from low-maternal education backgrounds have noisier responses, as reflected by greater spontaneous brainstem activity. Additionally, their response to the speech stimulus is less consistent over time with lower fidelity to the input signal.

**Conclusions:**
Noisier, more variable, and weaker brainstem responses are telltale signs of an inefficient auditory system. Consistent with work from laboratory animals, we interpret our findings as evidence that impoverished environments impinge upon the brainstem's ability to translate sounds into meaningful signals. We further suggest that disparate brain acuity may drive the achievement gap, effectively limiting social mobility for impoverished children. By studying SES within a neuroscientific framework, we have the potential to expand our understanding of how experience shapes the brain, in addition to informing intervention strategies.

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**Human brainstem plasticity: effects of auditory context and training**

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While it is well established that the auditory brainstem undergoes experience-dependent plasticity even in adulthood, mechanistic aspects of such plasticity are poorly understood in humans. Invasive studies on animal models show two forms of brainstem plasticity—a stimulus probability-based plasticity, and a second form of plasticity that is training-dependent. Whether these kinds of plasticity have manifestations in humans is still unanswered. Here we examine the frequency-following responses (FFR), which reflect ensemble neural phase-locking, to non-native pitch patterns using a passive oddball design. Participants listened to pitch patterns presented at a high probability of occurrence (standards) or a low probability of occurrence (deviants). They then underwent a 2-week sound-to-meaning training program during which they learned to use these pitch patterns to distinguish words. Post-training, the pitch patterns were presented again using the same passive oddball design. Results demonstrate the existence of two distinct forms of plasticity in the human auditory brainstem: a stimulus probability-driven plasticity that is present even before training and a training-dependent...
plasticity that alters the brainstem representation of pitch patterns. FFRs to ‘deviants’ were poorer than FFRs to ‘standards’ before training; post-training differences between standards and deviants are less salient. These results establish multiple, distinct forms of plasticity concurrently operational in the adult auditory brainstem.

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**Predictive coding in the human auditory brainstem: implications for neural plasticity**

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**Background:** A basic function of the human nervous system is to detect regularities in the sensory environment to predict future events. Neural models describing active prediction in the auditory system focus exclusively on cortical structures. Here we examined whether predictive coding is solely the domain of auditory cortical structures or if this coding represents a more systemic phenomenon that extends to subcortical structures.

**Methods:** We measured auditory brainstem potentials in adult subjects in response to an oddball paradigm in which the frequent stimulus was a complete and recognizable melody and infrequent stimuli consisted of the same melody with the final note either omitted or replaced with an unexpected note.

**Results:** Preliminary results show brainstem phase-locking for the final note of the melody for both the frequent and omitted stimulus conditions but not for a control condition. Moreover, the unexpected stimulus condition produces signals in the brainstem response at frequencies which are not present in the stimulus that correspond to the combination of the expected and unexpected notes.

**Conclusions:** Results indicate that anticipation of the final note of the melody is sufficient to drive phase-locked activity in the auditory brainstem, presumably through top-down, cortically mediated neural mechanisms.

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**Spatial coding, plasticity and functional asymmetries in the human auditory brainstem**

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The talk will explore interrelated themes of functional asymmetry, spatial coding, and plasticity in the human auditory brainstem. We report evidence for stimulus and context-dependent lateralization in the brainstem, measured with functional magnetic resonance imaging (fMRI). We found that speech, music, and noise stimuli are lateralized differently in the auditory thalamus. The lateralization of noise stimuli throughout the auditory pathway depends on the presence of other stimuli that are known to be preferentially processed in one hemisphere. A simple model in which the auditory cortices modulate their input from ipsi- and contralateral brainstem and thalamus via cortico-fugal projections accounts for these results.

We also present results on the encoding of interaural time differences (ITD) in the human brainstem. We presented binaural noises with different ITDs and recorded auditory brainstem revoked potentials using electroencephalography. We used an adaptation design to collect evidence for ITD encoding by two opponent neural populations, one tuned to the left and the other to the right acoustic hemifield. The results are consistent with models of ITD extraction in the auditory brainstem of other mammals. Lastly, we present results on plasticity in ITD coding in the brainstem from a study in which we shifted the correspondence between ITDs and sound locations by adding a constant time delay to one ear of test participants using small digital signal processing devices worn in the ear canal during several days. Most participants regained normal sound localization accuracy within 48 hours. ITD tuning in the cortex measured with fMRI changed in correspondence with the behavioral adaptation. Brainstem potentials recorded with EEG also showed a shift in directional tuning during the behavioral adaptation.

In summary, these results demonstrate an active role of the human auditory brainstem in perceptual and cognitive processes, such as context sensitivity, speech and music processing, and adaptation to modified sensory input.
Role of Cholinergic Modulation of the Auditory Cortex and Corticofugal Circuits in Spatial Learning in Adult Animals

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Background

Mammals are able to maintain a stable sound localization percept thanks to the ability of the brain to adapt to changes in the spatial cues available. We have shown that ferrets trained by positive conditioning to localize broadband sounds in azimuth are able to recalibrate their behavioral responses by re-weighting monaural and binaural cues altered after reversibly plugging one ear. Here we aim to establish the role of cortical acetylcholine (ACh) release in this training-dependent plasticity and to identify the neural circuits within the auditory pathway that are responsible.

Methods

We tested adaptation to a unilateral earplug in two different experimental conditions: (1) when cholinergic input to the auditory cortex was removed by injecting the immunotoxin ME20.4-SAP, which specifically targets the p75 neurotrophin membrane-bound receptor (p75NTR) expressed primarily by cholinergic neurons, in the nucleus basalis (NB) and (2) when pyramidal neurons from the primary auditory cortex that project to the inferior colliculus were selectively destroyed by chromophore-targeted laser photolysis.

Results

ME20.4-SAP injections in the NB produced a substantial loss of both p75NRT positive cells in this region and of acetylcholinesterase positive fibers throughout the auditory cortex. The loss of cholinergic neurons impaired the ability of ferrets to adapt to altered spatial cues following unilateral ear occlusion. After training they exhibited less complete recovery of localization performance than control cases, with adaptation taking place at a significantly slower rate. Although their ability to localize the long-duration sounds used to investigate plasticity was unaffected under normal hearing conditions, these animals had longer response times than controls. When the stimulus duration was reduced (<500 ms), they became less accurate than controls, with a direct correlation between the number of cholinergic cells lost and the impairment in performance.

When the projection from the primary auditory cortex to the inferior colliculus was selectively destroyed by chromophore-targeted laser photolysis in a second group of ferrets, the animals were again unable to adapt to relearn to localize sound after plugging one ear, but this time sound localization under normal hearing conditions was not affected.

Conclusions

Together, these results show that cortical cholinergic inputs contribute to auditory spatial perception and provide evidence for a role for ACh in adult auditory spatial learning. The cortico-collicular pathway is a key element in this adaptive plasticity, suggesting that collicular neurons can change their spatial response properties under cortical command.

The effects of music/language expertise on subcortical plasticity, auditory perceptual abilities, and cognitive transfer

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The scalp-recorded frequency-following response (FFR), an evoked potential with putative generators in rostral brainstem, provides a detailed window into subcortical encoding of complex sound including speech and music. Studies demonstrating enhanced FFRs in tone-language speakers and musicians suggest: 1) long-term experience with complex acoustic scenes strengthens brainstem response properties; 2) neural plasticity can emerge peripheral to cortex, with nascent effects observable even in subcortical auditory structures. Cross-domain comparisons show overall enhancements in FFRs elicited by either musical or linguistic stimuli in musicians and tone-language speakers alike. Thus, experience-dependent effects act to improve the brain’s initial registration of perceptually salient acoustic cues, a benefit which transfers across domains of expertise. Importantly, the fidelity (e.g., temporal precision and magnitude) of these sensory representations correspond with an individual’s degree of training/experience and, assuming the stimuli and task are behaviorally relevant to the listener, their performance in speech and music perception. These findings
collectively lead us to infer that both language and musical experience mutually benefit the subcortical extraction and subsequent perception of acoustic information. Yet, we also find that specific features of the auditory signal are highlighted in neurophysiological responses depending on their perceptual salience and function within a listener’s domain of expertise.

Alterations of the CIB2 calcium- and integrin-binding protein cause Usher syndrome type 1J and nonsyndromic deafness DFNB48

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Background
Many molecular components necessary for maintenance of vision and development of hearing have been discovered through identification of the genes underlying Usher syndrome. Usher syndrome (USH) is characterized by deafness and retinitis pigmentosa with variable age of onset and severity, sometimes accompanied by vestibular areflexia. We previously mapped a type I Usher syndrome locus segregating in two families and an autosomal recessive nonsyndromic hearing impairment (ARNSHI) locus (DFNB48) segregating in five families to chromosome 15q23-q25.1. Here, we report the identification of the gene causing USH1J/DFNB48 phenotype in humans.

Methods
We used genetic linkage analysis followed by a candidate gene screening strategy to identify the USH1J/DFNB48 gene, CIB2. Molecular modeling, biochemical, cellular, histochemical and physiological techniques in mice, zebrafish and Drosophila were used to determine the expression, localization and function of CIB2.

Results
We found that mutations in a calcium- and integrin-binding protein CIB2 are associated with nonsyndromic deafness DFNB48 and Usher syndrome type 1J (USH1J). CIB2 is a major cause of ARNSHI within the Pakistani population. In addition, a transition mutation of CIB2 co-segregated with ARNSHI in a Turkish DFNB48 family. In mice, CIB2 is localized in mechanosensory stereocilia of inner ear hair cells and in retinal photoreceptor and pigmented epithelium cells. Consistent with molecular modeling predictions, CIB2 significantly decreased the ATP-induced Ca\(^{2+}\) responses in heterologous cells, while DFNB48 alleles altered CIB2 effects on Ca\(^{2+}\) responses. Furthermore, in zebrafish and Drosophila, CIB2 is essential for the function and proper development of lateral line hair cells and retinal photoreceptor cells. We also show that CIB2 interacts with myosin VIIa and whirlin and thus is a new member of the vertebrate Usher interactome.

Conclusion
CIB2 mutations underlie Usher syndrome 1J and nonsyndromic deafness DFNB48. In Drosophila, CIB2 is critical for proper photoreceptor maintenance and function due to its involvement in calcium homeostasis. Since CIB2 is concentrated in stereocilia and interacts with myosin VIIa and whirlin, and Cib2 zebrafish morphants have reduced hair cell microphonic potential, we speculate that CIB2 regulate Ca\(^{2+}\) in hair cell stereocilia.
From Pattern Generator to Sound Receptor, Hair Cells Adjust Ca\textsuperscript{2+} Signaling to Their Function during Development

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Background
Ca\textsuperscript{2+}-triggered transmitter release from cochlear inner hair cells (IHCs) drives pre-sensory and sensory activity in spiral ganglion neurons (SGNs). Here, we studied the development of IHC Ca\textsuperscript{2+} signaling, afferent synaptic structure and SGN activity in mice.

Methods
We studied the changes of synaptic Ca\textsuperscript{2+} signal in mouse IHCs during development, using patch-clamp and fast confocal Ca\textsuperscript{2+}-imaging. Capacitance measurements and flash photolysis of caged calcium were used to investigate the apparent and intrinsic (biochemical) cooperativity of the hair cell ribbon synapse. Single unit recordings were performed in early postnatal mice to record activity of SGNs during the period of IHC maturation. Finally, immunofluorescence and electron microscopy were used to study molecular and morphological changes during development.

Results
Robust Ca\textsuperscript{2+} signals driven by spontaneous action potentials at IHC active zones (AZs) were observed before the onset of hearing. Concomitantly, we found bursting SGN activity \textit{in vivo}, which markedly differed from the Poisson-like spontaneous activity in hearing animals. At each synapse, several small appositions of AZs and postsynaptic densities (PSDs) were initially formed, in contrast to the single AZ/PSD pair present after maturation. Coupling between Ca\textsuperscript{2+} influx and vesicle fusion progressively tightened in postnatal IHCs (decrease in apparent cooperativity). In parallel, immunohistochemistry reported a progressive loss of extrasynaptic Ca\textsuperscript{2+} channels. In contrast to the decreasing whole-cell Ca\textsuperscript{2+} current and total channel number, synaptic Ca\textsuperscript{2+} signals increased in strength and heterogeneity, reflecting the emergence of AZs with more Ca\textsuperscript{2+} channels. Computational modeling suggested that these stronger AZs may drive the activity of SGN sub-population with low thresholds and high spontaneous rates.

Conclusion
Here, we characterized the synaptic Ca\textsuperscript{2+} signal associated with naturalistic electric activity of IHCs before and after the onset of hearing, namely, spontaneous action potentials and receptor potentials. We also demonstrated the presence of bursting activity in the postsynaptic SGNs \textit{in vivo}. The change in IHC function from spiking driver of pre-sensory activity to faithful encoder of sound intensity is accompanied by a confinement synaptic organization and function. Moreover, we propose that maturing IHCs heterogeneously distribute their Ca\textsuperscript{2+} influx among AZs, in order to decompose auditory information to functionally diverse SGNs and encode a broad dynamic range of sound pressure.
Mechanisms underlying heterogeneity of Ca$^{2+}$signaling among hair cell active zones

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Background

Sound intensity is encoded as the firing rates of spiral ganglion neurons. These neurons show different rate-level functions and are thought to collectively encode the large dynamic range of the auditory stimuli. We investigated the heterogeneous properties of presynaptic Ca$^{2+}$ microdomains and their relationship to synaptic ribbons. We tried to explain how hair cells decompose auditory information at their heterogeneous ribbon synapses driving neurons with different rate-level functions.

Methods

Postnatal (14-18 days) mouse inner hair cells from the apical half-turn of the organ of Corti were voltage clamped in whole-cell mode. A fluorescent Ca$^{2+}$ indicator (Fluo-8FF) and a ribbon reporter (TAMRA-conjugated CtBP2-binding peptide) were perfused into the hair cells via the pipette solution. Step or ramp depolarization protocols were used to evoke Ca$^{2+}$ influx and analyze the resulting Ca$^{2+}$ microdomains. A spinning disk confocal microscope was used to image each cell in 3D, and to characterize each Ca$^{2+}$ microdomain over time.

Results

Synaptic Ca$^{2+}$ microdomains and ribbons exhibited strong heterogeneity, and their fluorescence intensities were positively correlated. Additionally, we assessed the spatial distribution of ribbon and Ca$^{2+}$ hotspot intensities within individual inner hair cells. Larger ribbons and stronger Ca$^{2+}$ microdomains tended to localize to the modiolar (neural) side.

Conclusion

Our results imply that larger synaptic ribbons are associated with more Ca$^{2+}$ channels, which is expected to enhance neurotransmitter release at those synapses. Interestingly, previous studies (Merchan-Perez & Liberman, 1996) on cat cochlea suggested that high spontaneous rate auditory nerve fibers mainly innervate the pillar (abneural) face of inner hair cells. If conserved among species our finding of weaker Ca$^{2+}$ microdomains on the pillar face seems in conflict with this view. Further investigation is required to solve this apparent discrepancy.

Transmitter Release from Cochlear Hair Cells is Phase-Locked to Cyclic Stimuli of Different Intensities and Frequencies.

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Background

The auditory system analyzes time and intensity to code location of a sound source. These parameters are preferentially processed in different brainstem nuclei and so must be independently detected in the periphery and transmitted to the central nervous system. It is well-established that auditory nerve fibers are able to fire at a particular time within each cycle of a low-frequency periodic stimulus. Furthermore, this preferred phase can remain constant over a range of sound intensities at the characteristic frequency – phase constancy independent of intensity. We investigated this phenomenon at the first synapse of the hearing pathway, between inner hair cells and boutons of auditory nerve neurons.

Methods

Explants of the organ of Corti from neonatal rats were used for experiments (apical region). Simultaneous pre- and post-synaptic patch-clamp recordings at the inner hair cell-afferent synapse were performed. The series of differential equations corresponding to a model of the inner hair cell calcium sensor were solved numerically.

Results

Cyclic stimuli such as trains of depolarizations to different potentials were presynaptically applied, trying to mimic acoustic stimuli of different intensities. Synaptic responses elicited by these trains presented constant latencies (or phase) within each cycle, for a range of membrane potentials over which release probability varied significantly. Phase-locked release was observed with stimulating frequencies of 100, 250 and 500 Hz. In contrast, on single steps, synaptic responses proved to have a voltage-(calcium-) dependence on timing. Interestingly, short-term facilitation (only) partially compensated for changes in latencies (and also reduced jitter) in these single pulses. We also found that synaptic depression elicited longer 1st latencies events. Thus, during repetitive presynaptic stimulation, as hair cells are more strongly depolarized, increased calcium channel gating would speed up transmitter release, but the resulting vesicular
depletion produced a compensatory slowing. Quantitative simulation of ribbon function showed that these two factors will vary reciprocally with hair cell depolarization (stimulus intensity) to produce constant synaptic phase.

**Conclusion**

We propose that equilibrium would be established between each stimuli level and the degree of synaptic depression. Stronger stimuli accelerate release kinetics but operate on a smaller vesicular pool, which in contrast, slows down the release. Weaker pulses produce less depression determining a larger pool, which in turn, favors faster kinetics. This phenomenon helps determining that in trains of stimuli the average phase is conserved.

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**Otoferlin Regulates Hair-Cell Receptoneural Secretion Through Novel Protein Interactions**

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**Background**

Emerging evidence supports a pivotal role for otoferlin in the regulation of hair-cell secretion/exocytosis, where this protein may sense calcium in the process of SNARE-driven vesicle fusion.

**Methods**

These studies involved either purified, bacterially-expressed C2 domains and SNARE motifs, or full-length proteins as expressed in rat brain. In vitro binding was measured via surface plasmon resonance analysis, pull-down assays, and immunoprecipitation. Calcium binding of otoferlin C2 domains was detected by isothermal calorimetry. Proteomic analysis using mass spectrometry was carried out on immunoprecipitated proteins. Immunolabeling of rat organ of Corti was performed with anti-otoferlin and anti-Ca₉.1.3 antibodies.

**Results**

Calorimetric measurements demonstrated Ca²⁺ binding to the otoferlin C2F domain. MS-proteomic analysis of otoferlin immunoprecipitation products showed that syntaxin 1B and SNAP-25 specifically interact with otoferlin. RT-PCR revealed expression of syntaxin 1B in rat organ of Corti. In vitro analysis by SPR and GST pull-down indicated that the SNARE motif of syntaxin 1 specifically interacts with otoferlin C2 domains, enhanced by addition of Ca²⁺ up to 200 µM. C2D, C2E, and C2F showed similar association-binding kinetics with the SNARE motif for increasing levels of Ca²⁺, but the dissociation rate at 20 µM Ca²⁺ was significantly reduced compared to dissociation rates at 0 µM and 10 µM Ca²⁺. There was no substantial further decrease in dissociation rate with Ca²⁺ increments up to 200 µM. We also demonstrated, for the first time, that C2 domains of otoferlin bind each other in vitro in the absence of Ca²⁺. Upon addition of Ca²⁺ to the binding buffer, this interaction was substantially diminished.

**Conclusion**

Our results indicate that otoferlin may be an unusual calcium sensor, playing an important role in vesicle fusion in hair cells. The otoferlin C2F domain binds calcium, and interacts with the syntaxin 1 SNARE motif in a calcium-dependent manner. Otoferlin selectively interacts with syntaxin 1B, which is a syntaxin form present in rat organ of Corti. The otoferlin C2-syntaxin 1B SNARE interaction is possibly stabilized at 20 µM Ca²⁺ due to decreased dissociation at that Ca²⁺ level. Our finding that C2 domains engage in mutual C2-domain binding in the absence of calcium indicates a possible intramolecular mechanism involving closed otoferlin at low Ca²⁺ and open otoferlin at higher Ca²⁺ that may, in turn, regulate otoferlin’s intermolecular interactions upon synaptic influx of Ca²⁺.
How Different Human Electrophysiology and Neuroimaging Methods Can Help Us Understand Auditory Processing
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Background
As organizers of this Young Investigator Symposium, we will use this talk to introduce the multiple methods described in this symposium’s presentations to junior scientists as well as the interested but uninitiated.

Methods
We will discuss how non-invasive electrophysiological and neuroimaging methods (fMRI, ABR, EEG, and MEG) work, what types of experimental designs are used in conjunction with them, and how their varying spatial and temporal resolutions offer complementary information.

Results
We will provide a brief introduction to the talks in this symposium, highlighting different labs’ approaches to answering questions of common interest.

Conclusion
Electrophysiology and neuroimaging, in concert with recent developments in data analysis methodology, provide new insights into how humans listen.
Bottom-Up and Top-Down Contributions to Individual Differences in Auditory Spatial Attention Task Performance
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Background
The ability to attend to a sound of interest amidst competing sounds is vital for navigating through the complex auditory scenes of everyday life. Even among “normal hearing” listeners, there is considerable variability in auditory spatial attention task performance. Our previous findings indicated that fidelity of brainstem phase locking to simple stimuli as measured using the brainstem frequency following response (FFR) is predictive of spatial attention task performance. Here, we extend the study of bottom-up coding using FFRRs to understand the effect of sound level on envelope phase locking up to the brainstem, and use Magnetoencephalography (MEG) to study oscillatory synchronization in the cortex during top-down selective attention preparation.

Methods
MEG was recorded from normal-hearing listeners presented with three simultaneous streams of spoken letters, from distinct spatial locations using inter-aural time disparities (ITDs) or binaural room impulse responses (BRIRs). Subjects were visually cued to attend to either the left or the right location and count the occurrences of a target letter. MRI scans provided anatomical constraints for MEG source analysis. In a separate session, FFRRs to a 100 Hz click-train at 3 different levels (60, 70 and 80 dB SPL) and both polarities were recorded using a 32 channel EEG cap. After optimally phase aligning the FFRRs across channels, envelope phase-locking values (PLVs) were computed using a bootstrap procedure.

Results
FFR results indicate that PLVEnv consistently increases when stimulus intensity increases from 60 to 70 dB SPL. However, there are large individual differences in the change in PLVEnv from 70 to 80 dB SPL that correlate with performance on the spatial attention task. Preliminary MEG findings show subject-specific increase in beta-band activity in the right Intraparietal Sulcus (rIPS) in the preparatory period (post cue, before sound) in the attend trials and particularly in trials where the attended location is switched from the previous trial.

Conclusions
Subtle supra-threshold envelope spatial cues are important particularly in crowded and reverberant settings. Robustness of envelope coding at high levels, which can be investigated at the individual level using FFRRs, may reflect the fidelity of integration of information from low and high threshold auditory nerve fibers. Sub-clinical effects of noise-exposure, aging etc. that compromise this may hence be objectively assessed. MEG correlates of top-down spatial selective attention along with FFRRs help explain the large individual variability in spatial attention performance.

Neuromagnetic signatures of segregation in complex acoustic scenes
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The natural auditory environment consists of multiple dynamically varying sound sources. In order to make sense of this complex mixture of sounds, we need to segregate individual sound sources. The brain has evolved specialized mechanisms for analyzing auditory scenes but the underlying mechanisms remain to be fully explained.

We improved upon earlier paradigms based on deterministic patterns of pure tones and modelled acoustic scenes using a stochastic figure-ground stimulus (SFG, Teki et al., 2011). The stimulus comprises a series of chords (25 ms long) containing random frequencies that vary from one chord to another in a range from 200 Hz to 2.5 kHz. To study segregation, we introduced a figure by randomly selecting a certain number of frequencies (‘coherence’) and repeating them over a certain number of chords. This manipulation allows us to parametrically control the salience of the figure, which can only be extracted by binding across both time and frequency. We found that listeners are very sensitive to the emergence of these complex figures. We previously established a role for the intraparietal sulcus in stimulus-driven segregation of these figures (Teki et al., 2011) and ongoing work suggests a role for temporal coherence in mediating segregation (Shamma et al., 2011).

We used MEG to investigate mechanisms underlying the emergence of figures with different salience (coherence of 2, 4 or 8: 0.6s long) presented after statistically similar background segments (0.6 s). Listeners were engaged in an incidental visual task and were naive to the manipulations within the SFG stimulus. In another condition, we presented the same
Analysis of time-locked activity in the auditory cortex shows an initial onset response to the emergence of the figure, followed by a sustained response which follows the figure. Source analysis is ongoing. The figure onset responses occur about 100 ms in the coherence=8 condition, and 150 ms for coherence values of 4 and 2. These latencies correspond to a duration of 4 (or 6) chords and parallel behavioural performance (obtained separately). Latencies from the ‘noise’ condition reveal the same threshold, suggesting that the segregation mechanism was not affected by the interspersed noise bursts. Time-frequency analysis is ongoing using a beamformer approach (Sedley et al., 2012) to identify sources of early and late oscillatory activity.

**Cortical Encoding of Speech in Challenging Listening Environments**

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Normal hearing listeners are superb at understanding speech in noise. Here, we study the neural underpinnings of this phenomenon in human auditory cortex, by recording from subjects listening to spoken narratives using magnetoencephalography (MEG). In one experiment, the spoken narrative is mixed with spectrally matched stationary noise at different signal to noise ratios (SNR). We analyzed the auditory cortical response synchronized to the slow temporal modulations of speech (< 10 Hz). It is found that the cortical synchronization to the very slow temporal modulations of speech (< 4 Hz) is stable until the noise is far stronger than the speech. The cortical synchronization to faster modulations (> 4 Hz), however, is more susceptible to noise. In another experiment, the stimuli are processed by noise vocoders, to simulate listening with cochlear implants. The neural synchronization to vocoded speech is greatly reduced even when the background noise is only mildly weaker than speech, consistent with the behaviorally observed noise susceptibility of vocoded speech. Taken together, these results suggest that the neural representation of very slow temporal modulations of speech is very robust to noise. This robustness, however, relies on the spectro-temporal fine structure of speech.

**Using high-field fMRI and multivariate methods to study neural representations of complex sounds in human auditory cortex**

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**Background:**
Investigating auditory processing in the human brain remains a challenge due to the small size of auditory cortex, substantial individual differences in anatomy, and limitations of the methods for neural measurement. We used functional magnetic resonance imaging (fMRI) in combination with multivariate analyses methods to assess which acoustic and abstracted features of complex natural sounds are encoded in human auditory cortex during perception and cognitive tasks such as change detection and imagery.

**Methods:**
Structural and functional images were acquired on a 3T Siemens Tim Trio and a 7T Agilent/Siemens MRI scanner. We evaluated the benefits and drawbacks of ultra-high field 7T MRI to assess auditory cortex function at higher spatial resolution, used new acquisition sequences that reduce scanner noise and applied multivariate methods that are insensitive to individual differences in anatomy to make fMRI more suitable for auditory research. Participants listened to, performed change detection on and imagined simple and complex, natural sounds. Multivariate pattern analysis (MVPA) was conducted to assess feature tuning in human auditory cortex and to determine whether the acoustic and abstracted features of sounds encoded were task-dependent.

**Results:**
Our results show that the information encoded in spatially distributed patterns of activity in auditory cortex differs depending on the cognitive task being performed. During perception and short-term memory maintenance activity patterns in auditory cortex contained stimulus-specific information while abstracted, categorical information was encoded in the same region during auditory imagery. Furthermore, differences in activation magnitude and pattern distinctiveness were related to individuals’ memory capacity and perceived imagery vividness. This indicates that auditory cortex is recruited for processes beyond analyzing simple feature information, playing an important role in maintaining sustained representations of sounds in short-term memory and encoding abstracted information during imagery. A comparison of
data acquired on the 3T Siemens Tim Trio and 7T Agilent/Siemens MRI scanner additionally indicates that ultra high-field MRI is particularly well suited for the assessment of human auditory cortex.

Conclusions:
Ultra-high field fMRI and multivariate methods are well suited for analyzing feature coding in human auditory cortex. They allow for the identification of much finer detail both structurally and functionally. In combination, these methods provide exciting opportunities to investigate how neural representations in auditory cortex vary within as well as between individuals.

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What Musicians Reveal About the Aging Brain: Behavioral and Neural Indices
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Background: As the world’s population ages, there is increased interest in identifying lifestyle factors that reduce aging’s deteriorating effects. One marker of healthy aging is the ability to remain socially active, which requires intact communication skills. A major complaint of older adults, even those with normal hearing, is the inability to understand speech, especially in the presence of background noise. Defining the effects of age on auditory neurobiological processes is important as we seek to remediate this deficit. On a neural level, aging is associated with reduced temporal synchrony, evidenced by delayed neural timing and increased trial-by-trial response variability. Given recent evidence that older musicians have enhanced speech perception in noise, we asked whether musical experience offsets age-related declines for the neural processing of speech.

Methods: We recorded neural responses to a speech sound /da/ in 140 normal hearing younger (18-30) and older (45-65) normal hearing musicians and nonmusicians matched for age, IQ and sex, focusing on neural timing and response variability. Furthermore, we quantified both subjective and objective indices of speech-in-noise perception by administering a self-report questionnaire on which each individual rated themselves on their ability to hear in noise and the Hearing in Noise Test.

Results: Older nonmusicians have delayed neural responses and increased response variability relative to older musicians, who demonstrate resilience to these effects of aging and perform similarly to younger adults. Older musicians reported higher ratings on the scale of self-perceived hearing-in-noise ability, which was confirmed by better speech-in-noise perceptual thresholds, suggesting fewer communication difficulties relative to older nonmusicians.

Conclusions: Musical training strengthens aspects of neural processing that degrade with age, which may account for their enhanced ability to hear in noise. As such, we propose that older musicians may serve as a model of plasticity which can be used to further our understanding of the aging process on the nervous system.

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Hierarchical neurocomputations underlying concurrent sound segregation: Connecting periphery to percept
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A critical process in auditory scene analysis is the ability to perceptually separate concurrently occurring sounds. When acoustic energy is harmonically related across the frequency spectrum, an auditory event is perceived as a single sound object; when there is additional energy not related to the same f0, listeners often report hearing two sound objects. Numerous studies have found that in human listeners, the perception of concurrent sounds is paralleled by complementary changes in cortical event-related potentials (ERPs). Although these studies provide a window into the neural mechanisms governing concurrent sound segregation, the neural architecture and hierarchy of neurocomputations supporting this robust perceptual process remain poorly understood. Here, we employ a coordinated blend of techniques including neurocomputational modeling, cortical/subcortical ERPs, and human psychophysics to assess the role of both peripheral and central neurophysiological responses in formulating concurrent sound segregation. Results suggest that the acoustic cues necessary for segregating auditory objects (e.g., inharmonicity) are adequately coded at the level of
auditory nerve, are maintained with high fidelity within response properties of the upper brainstem, and are further transformed into a perceptual correlate by early auditory cortex. Our novel approach in studying multiple brain indices simultaneously helps illuminate the neural mechanisms underlying concurrent sound segregation and may lead to the further development and refinement of physiologically driven models of auditory scene analysis.

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**Investigating BOLD fMRI tuning to interaural level and time differences in human auditory cortex**

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**Background:** Interaural time and level differences (ITD and ILD) are binaural cues important in sound localization. At present, mechanisms underlying their cortical representation remain unclear. Responses from binaurally-tuned neurons suggest that a majority in each hemisphere respond more to earlier and more intense stimulation of the contralateral rather than ipsilateral ear. However, evidence for contralateral tuning in blood oxygenation level-dependent (BOLD) fMRI of human auditory cortex (hAC) is more equivocal, possibly due to the nature of the measure, stimuli presented, and/or potential ‘carryover’ effects of stimulus history. The present study will therefore investigate binaural processing in hAC by measuring direct and carryover effects of parametric binaural cue modulation on the BOLD response.

**Methods:** In a series of experiments, the BOLD signal was measured in response to modulation of (Exp. 1) ILD (0, ±5, ±10, ±20, ±30dB) and (Exp. 2) ITD (0, ±200, ±500, ±800, ±1500μs) imposed on narrowband click trains (4000 Hz carrier frequency, 1.8 kHz half-max bandwidth, 500 Hz click rate). Exp. 3 employed identical ITDs imposed on broadband stimuli (trains of 1ms Gaussian noise bursts repeating at 100 Hz). Across experiments, stimulus trains were presented for 1s (interstimulus intervals jittered from 1-5s), at 80 dB SPL, via insert earphones (Sensimetrics). Binaural cue order was counterbalanced, following a continuous carryover design [Aguirre, NeuroImage 35:1480-94, 2007]. The task involved pitch detection orthogonal to the experimental manipulation. Continuous event-related images were acquired (TR=2s, 42 slices, 2.75 x 2.75 x 3mm resolution, 3T). Data pre-processing and individual analyses were conducted using FSL; cross-subject analyses were conducted on the cortical surface using Freesurfer and Matlab.

**Results:** Consistent with the neural response data, a contralaterally-tuned non-monotonic ILD-response function was observed, with maximal responses at extreme contralateral ILDs. Responses declined at moderate contralateral and ipsilateral values but increased again at extreme ipsilateral ILDs. In contrast, ITD-response functions for both narrow and broadband sounds showed little to no contralateral tuning. Stimulus history effects also differed across cues, with preliminary ILD results suggesting attenuation of the response to ipsilateral stimulation with ipsilateral stimulus repetition.

**Conclusions:** Findings will be discussed in the context of binaural processing models, such as coding by excitatory/inhibitory “opponent” populations, by temporal spike patterns, and by distributed codes across populations of panoramic neurons. [Supported by R01-DC011548, R03-DC009482.]
Using fMRI to study the role of attention in speech perception.

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Background

The conditions of everyday life are such that people often hear speech that has been degraded (e.g., by background noise or electronic transmission) or when they are distracted by other tasks. However, it remains unclear what role attention plays in processing speech that is difficult to understand. Functional magnetic resonance imaging (fMRI) provides a valuable method to explore this issue by allowing the comparison of brain responses to stimuli that are attended and unattended. In the current study, we used fMRI to assess the degree to which spoken sentences were processed under distraction, and whether this depended on the acoustic quality (i.e., the intelligibility) of the speech.

Methods

On every trial, participants were cued to attend to one of three simultaneously presented stimuli: a spoken sentence, an auditory distracter (a sequence of noise bursts that did not interfere with the speech signal), or a visual distracter (a series of rotating ellipses). Regardless of which stimulus was attended, the sentence on every trial was presented at one of four levels of intelligibility: clear speech, high- or low-intelligibility noise-vocoded speech, or rotated noise-vocoded (i.e., unintelligible) speech. After scanning, a surprise recognition test was administered to measure subjects' memory for the sentences.

Results

The post-test showed that attention did not affect the recognition of sentences presented in a clear or unintelligible form: clear sentences were always recognized at better than chance levels, regardless of whether or not they had been attended, and unintelligible sentences were never recognized. However, attention significantly enhanced the recognition of high- and low-intelligibility noise-vocoded speech.

Furthermore, the fMRI results demonstrated that speech-sensitive cortex could be parcellated according to how speech-evoked responses were modulated by attention. Responses in auditory cortex and areas along the superior temporal sulcus (STS) took the same form regardless of attention, although responses to distorted speech in portions of both posterior and anterior STS were enhanced under directed attention. In contrast, frontal regions, including left inferior frontal gyrus, were only engaged when listeners were attending to speech and these regions exhibited elevated responses to degraded, compared with clear, speech.

Conclusions

We suggest this pattern of fMRI responses is a neural marker of effortful listening. Together, our results suggest that attention enhances the processing of degraded speech by engaging frontal-cortex mediated mechanisms that modulate perceptual processing of speech.
Stereocilia deflection phase is frequency dependent
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Background
Mammalian hearing relies on the movements of cochlear stereocilia. Thanks to their attachment to the tectorial membrane (TM), outer hair cell (OHC) stereocilia bundles are deflected by sound-induced shearing motion between the TM and the cuticular plate. There is a consensus that such is not the case for the freestanding inner hair cell (IHC) stereocilia bundles whose deflection is driven by sound-induced viscous drag of the endolymph. Following these differences in stimulatory forces, the receptor potential recorded in IHCs shows a phase lead of 90° relative to the basilar membrane motion. If the OHC stereocilia deflection depends only on the basilar membrane movement as generally assumed, then the stereocilia deflection in IHC would also have a 90° phase lead relative to OHC stereocilia. To investigate this hypothesis, direct measurement of the stereocilia deflection phase is required.

Methods
Using temporal bone preparations from guinea pigs, we measured the deflection phase at different frequencies and sound stimulus intensities by time-resolved confocal imaging and optical flow algorithms, a technical approach that we recently used to characterize nanometre-scale sound-evoked length changes in OHC stereocilia.

Results
IHC stereocilia bundles with respect to OHC stereocilia bundles exhibited a phase lead of about 60°, smaller than expected. In addition, this phase difference was maintained across a range frequencies and intensities. In both IHCs and OHCs, the phase increased with frequency but not with the stimulus intensity.

Conclusion
Taken together, these observations suggest that the mechanism for driving IHC bundles may be more complex than previously envisioned.

Synchronization of active hair bundles by artificial membranes
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Background
In vitro, mechanically decoupled or free-standing hair bundles from the frog sacculus exhibit spontaneous mechanical oscillations. These active hair bundles show signs analogous to the “active process” seen in vivo in the auditory systems of all vertebrates, suggesting that hair bundle motility plays a role in active amplification. However, the spontaneous oscillations are uncorrelated and the performance of a single bundle does not match that of the auditory system in vivo. Several theoretical studies have suggested that mechanical coupling of active hair bundles can lead to an enhancement of the single-cell active process. We used artificial membranes to couple a small number of active hair bundles and recorded both spontaneous and entrained movements.

Methods
We used an in vitro preparation of the frog sacculus in which the sensory epithelium is mounted in a two-compartment chamber mimicking the native ionic environment. Artificial membranes were fabricated from poly(lactic-co-glycolic acid) (PLGA) microspheres. The microspheres were briefly melted to yield approximately hemispherical shapes, and the artificial membranes were coated with concanavalin A to improve adhesion to the hair bundles. The artificial membranes were dispersed onto the preparation and typically coupled two to four hair bundles. In several experiments, the artificial membranes were mechanically stimulated with flexible glass fibers driven by a piezoelectric stack actuator. Bundle and artificial membrane motion were recorded with a high-speed CMOS camera as described previously.

Results
When coupled by artificial membranes, active hair bundles exhibited robust spontaneous oscillations. The motion of the hair bundles was strongly correlated (correlation coefficients of about 0.9) suggesting that direct mechanical coupling can lead to synchronized motion of a small number of active hair bundles.

Conclusion
When mechanically coupled, active hair bundles can exhibit synchronized spontaneous oscillations. The forces generated by a few hair bundles are sufficient to move overlying accessory structures. Preliminary data suggest that when driven
sinusoidally, the bundles in this coupled system can exhibit sharpened frequency tuning relative to a single free-standing bundle. Further experiments on this hybrid system are ongoing.

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Cable property of OHC lateral membrane
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Background
The outer hair cell’s (OHC) fast mechanical response is capable of following sinusoidal stimuli that go beyond 100 kHz. It remains puzzling that such a mechanical response can be achieved with the lateral membrane as low-pass filter. We hypothesize that the subsurface cisternae serve to overcome the restraining effect of lateral membrane’s low-pass filter function therefore allowing the OHC’s electromotility to respond to fast stimuli. We measured membrane time constants at various locations along the OHC lateral membrane by using a two electrode technique.

Methods
Isolated guinea pig OHCs are whole cell recorded with a perforated patch electrode near the supra-nucleus region. An electrical stimulus was generated by a roving electrode along the lateral membrane. Recordings were made with series resistance compensated to 10 Mohm under all conditions. Whole cell condition was then established and measurement repeated. Measurements were also made after OHCs were treated with 10 mM salicylate for 40 minutes. Membrane time constants were compared.

Results
The time constant for different positions along the lateral membrane showed little change when the OHC plasma membrane was intact (perforated patch) but showed a distance dependent change (re: stimulating electrode) when the membrane was ruptured (whole cell). We believe that the sub-plasma membrane space is destroyed in this case. Such changes can be reproduced by treating OHC with 10 mM salicylate even during perforated patch recording. Salicylate can also destroy the OHC’s sub-plasma membrane space.

Conclusion
The intact subsurface cisternae helps to overcome the membrane filter that allows the OHC to respond to high frequency transduction current.
Optimal Electrical Properties of Outer Hair Cell Membrane for Cochlear Amplification

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Background
The organ of Corti (OC) is an auditory epithelium specific to the mammalian cochlea comprising sensory hair cells and supporting cells riding on the basilar membrane. The outer hair cells (OHCs) are cellular actuators that amplify small sound-induced vibrations for transmission to the inner hair cells. Two recent observations provide important insights to current debate on cochlear amplification by OHCs. First, OC with active OHCs was observed to vibrate with a characteristic mode such that the top and bottom surface of the OC have 90 degree phase difference (Chen, Zha et al., 2011). Secondly, the conductance of OHC membrane varies along the cochlear coil such that it is higher at high frequency location (Johnson, Beurg et al., 2011). We investigated the implications of these two findings through computational analyses of the OC.

Methods
Developed finite element model of the OC incorporates the sophisticated OC geometry such as pillar cells, tilted Deiters cell processes and OHCs, and longitudinal coupling (Nam & Fettiplace, 2010). The model also incorporates force generation by OHCs originating from active hair bundle motion due to gating of the transducer channels and somatic contractility due to the membrane protein prestin. It was a time-domain analysis with fully non-linear OHC mechano-transduction and electro-motility.

Results
The 90 degree phase difference between the top and the bottom surface vibrations of the OC of appeared only when the OHC motility is prominent. Simulations of an individual OHC show that the OHC somatic motility lags the hair bundle displacement by ~90 degrees. Prestin-driven contractions of the OHCs cause the top and bottom surfaces of the OC to move in opposite directions. Combined with the OC mechanics, this results in ~90 degrees phase difference between the OC top and bottom surface vibration. An appropriate electrical time constant for the OHC membrane was required to achieve the phase relationship between OC vibrations and OHC actuations. When the OHC membrane conductance is too high or too low, the OHCs do not exert force with the correct phase to the OC mechanics so that they cannot amplify. This optimal OHC membrane conductance is in nice agreement with the measured value of Johnson et al.’s (2011).

Conclusion
There exists an optimal range of OHC electrical property to set the correct phase relations needed for cochlear amplification.
Meno presto, prestin: disparities between voltage-sensor charge and electromotility reveal slow chloride-dependent molecular state transitions in SLC26a5
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¹Yale University

Background
Outer hair cells (OHC) drive cochlear amplification, which enhances auditory detection and discrimination. The motor protein, prestin, which evolved from the SLC26 anion transporter family, underlies the OHC’s electromotility.

Methods
Voltage clamp and video analysis of OHC NLC and motility.

Results
Here we report on simultaneous measures of prestin’s voltage sensor charge movement (NLC) and electromotility that evidence disparities in their voltage dependence and magnitude as a function of intracellular chloride. A simple kinetic model, possessing fast anion binding transitions and fast voltage dependent transitions, coupled together by a much slower transition recapitulates these disparities and a host of other biophysical observations on the OHC. The intermediary slow transition probably relates to the transporter legacy of prestin, and this intermediary gateway which shuttles anion bound molecules into the voltage-enabled pool of motors, provides molecular delays that present as phase lags between membrane voltage and electromotility. Cochlear modelers find that phase lags are required to effectively inject energy at the appropriate moment to enhance basilar membrane motion.

Conclusion
Thus, slowing down prestin has its benefits.

Remodeling the intracellular distribution of DNA repair enzymes in spiral ganglion neurons in response to noise stress and otoprotective therapy
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Background
Intracellular stress gradients drive the spatial distribution of DNA repair enzymes. In this study, we investigated if spiral ganglion neurons could remodel intracellular distributions of DNA repair enzymes in response to stress and otoprotective therapy.

Methods
The intracellular locations of DNA excision repair enzymes were determined within spiral ganglion neurons under normal conditions, after noise exposure, and following otoprotective treatment with carboxy alkyl esters (CAEs). Line scans were used to record immunoreactivity within the soma of each neuron to objectively profile the subcellular distribution of the enzymes.

Results
After noise stress at 105 dB for 4 hours, the enzymes aggregated in the cytoplasm and this effect was associated with a significant loss (p < 0.001) of neural sensitivity measured by compound action potential. After CAE therapy (160 mg/kg/28 days), the enzymes were enriched within multiple intracellular locations and this response had no significant change in neural sensitivity (p > 0.05). CAE therapy provided to noise exposed animals also resulted in the enrichment of repair enzymes in multiple intracellular compartments. This response was associated with significant preservation of neural sensitivity (p < 0.05).

Conclusion
In summary, the current study revealed that spiral ganglion neurons exhibit multiple compartmentalizing modes for excision repair enzymes and these modes exhibit plasticity following noise stress and otoprotective treatment. Furthermore, regulation of intracellular gradients of DNA excision repair enzymes may represent a novel approach to preserving neural function following stress.
Genetic/transgenic conditional expression of nonmuscle myosin-II in auditory neurons: Head domain regulates assembly of the α-helical coiled-coil tail
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Background
Nonmuscle myosin-II (MyoII) is an actin-binding protein that is involved in several motor functions including growth cone motility and neural migration, outgrowth and retraction. The molecule can be functionally divided into a head-domain and a α-helical coiled-coil tail. Self-assembly of the MyoII α-helical coiled-coil tail with other MyoII molecules form bipolar filaments that power cell movements. This self-assembly of the tail is intrinsically regulated by intramolecular interactions of the tail with the head-domain as demonstrated by in vitro experiments. However, there is no in vivo evidence for this regulatory mechanism.

Methods
In this study, we further define whether the head-domain of MyoII regulates in vivo self-assembly of the α-helical coiled-coil tail in auditory neurons. A GAL4-UAS gene expression system was used to selectively express zip/MyoII full-length, zip/MyoII tail with and without isoleucine-glutamine (IQ) motifs in Drosophila melanogaster auditory (Johnston's organ) sensory neurons. The N-terminus of each construct was fused to green fluorescent protein (GFP) to follow the distributions of transgene expression.

Results
Our results reveal that the full-length molecule exhibited either globular or rod conformations in auditory neurons of Johnston's organ. In the absence of the head domain, the tail self-assembles into filament-like rods in both fixed and living preparations of Johnston's organ neurons. The tail+IQ motif failed to form rod conformations and instead filled the neuronal cell bodies with diffuse fluorescence.

Conclusion
The results suggest that the head-domain is critical for modulating the self-assembly of the tail in vivo. These findings may help to explain why human mutations in the head domain result in severe diseased phenotypes.
**Are Species Differences in Dip Listening Predicted by Species Differences in Temporal Processing? An Evoked Potential Study in Treefrogs**

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**Background**

Acoustic communication in large social groups is made difficult by high noise levels, which can mask signals of interest. One solution for coping with this problem involves listening in the “dips” of fluctuating background noise, which requires the ability to track temporal envelope fluctuations. Acoustic communication in social groups is not a uniquely human problem, nor is dip listening a uniquely human solution. We have previously found that one species of treefrog, Cope’s gray treefrog (*Hyla chrysoscelis*), is also able to exploit dips in background noise to recognize communication signals at lower thresholds, while a related species, the green treefrog (*Hyla cinerea*), appears incapable of doing so. Here, we tested the hypothesis that these species differences in dip listening are due to species differences in temporal processing.

**Methods**

To test our hypothesis, we used two measures of evoked potentials to assess temporal processing in gray treefrogs and green treefrogs. For the first measure, evoked potentials were recorded in response to double-click stimuli. Temporal integration time corresponds to the shortest inter-click interval at which distinct responses are evoked by both clicks. The second type of evoked potential, termed the auditory steady-state response (ASSR), measured phase locking by the auditory nerve to the envelope of amplitude-modulated tones. The magnitude of the fundamental frequency of the ASSR was used to construct modulation rate transfer functions (MRTF).

**Results**

Assessments of temporal processing using these two methods revealed very small differences between the two species. In both species, we observed two distinct responses to clicks with inter-click intervals averaging as low as 3-4 ms. The average inter-click interval at which two responses were present was slightly shorter in gray treefrogs than green treefrogs. The MRTFs for the two species were broadly similar in shape and did not differ greatly in absolute magnitude.

**Conclusion**

Our results indicate that temporal processing abilities at the level of the eighth nerve are conserved between the two species. While small species differences were observed, they are probably not great enough to account for species differences in dip listening. Our data do not rule out species differences in temporal processing at higher levels of the ascending auditory system. [This work was supported by NSF IOS 0842759.]

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**Encoding temporal fine-structure at different sound levels in the barn owl auditory nerve**

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**Background**

Barn owls use interaural time different (ITD) for sound localization. In order to do so their auditory system encodes the fine structure of sound up to 10 kHz. The level of ecologically relevant sounds can vary by tens of decibels and the effect on the monaural encoding of sound could have a great effect on ITD tuning. It is therefore important to understand time encoding at the first stage of the auditory pathway at different sound levels.

**Methods**

We recorded responses to noise of auditory nerve fibers with best frequency (BF) spanning the entire audible frequencies of owls. Unfrozen noise at three intensities was presented: one slightly above threshold, one in the middle of the dynamic range, and one at saturation. To compute the AN impulse response, we used reverse correlation on the response to unfrozen noise, the spike triggered averaged (STA). Resulting STA were fitted to linear gamma chirps.

**Results**

While some properties of impulse responses were in concordance with studies in mammals and in the barn owl, some important differences were noticed. Unlike in mammals, the impulse responses showed frequency glides for fibers with BFs above 4 kHz. The glides were positively correlated with BF and had larger magnitude than previously measured in downstream nuclei. For lower BF the glide slopes were very small and independent of frequency. More importantly, the glide slopes were significantly dependent on level for BFs above 5 kHz. In this high frequency range the changes in slope could be as big as 1kHz/ms when level was varied across the dynamic range. In addition, the sign of the slope could
change. We show that the outputs of the AN filters at different levels did not only have different envelopes but also different fine structures. Using cross-correlation of the analog filter outputs we predicted that the shape of the ITD curves derived from those neurons would be greatly dependent on interaural level difference (ILD).

Conclusion
While invariance on ILD would be desired to encode ITD our data suggest that, for BFs>5 kHz, the fine structure of the responses at the first level of the monaural pathway is not level invariant. This casts additional constrains on the downstream pathway, in order to encode ITD independently of sound level, as it has been shown. More generally, level dependence of the spike train fine structure could be functionally useful in tasks such as distance or loudness perception.

Development of Spiral Ganglion Neuron Intrinsic Excitability and Modulation by NT-3 in CBA Mice
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Background
The neurotrophins BDNF and NT3 have critical roles in promoting the survival, connectivity and function of spiral ganglion neurons (SGNs). In 1 week old mice the features of SGN intrinsic excitability are tonotopically organized with rapidly accommodating, high threshold neurons found in the base and slowly accommodating, low threshold neurons located in the mid-apical ganglion (Liu & Davis, J. Neurophysiol. 2007). Here we assess timing and excitability features including threshold, accommodation and action potential latency from P1 to P14 and have begun to determine the role of NT3 application in modulating these features during development.

Methods
Whole-cell current-clamp recordings were performed on SGN cultures obtained from postnatal day 1-14 (P1-P14) mice to ascertain the features described above. NT-3 (10 ng/ml) or vehicle was applied at time of plating. Step current injections (240 ms duration) were delivered to assess voltage threshold, action potential (AP) latency and accommodation.

Results
From P1 to P7, we find that base and apex neurons retain the aforementioned tonotopic profile for P6 and P7, with P5 being a transition period. However, base neurons from P1 to P4 manifest a prominently more excitable and immature phenotype relative to P7 base neurons: they display lower thresholds (P7 base vs. P4 base) -42.5±0.9 mV, n=25 vs. -48.3±1.1 mV, n=15; P<0.01), longer latencies (7.4±0.4 ms vs. 15.8±0.8 ms; P<0.01) and slow accommodation (1.24±0.13 vs. 13.2±2.7; P<0.01). From P7 to P14, both apex and base neurons become faster and rapidly accommodating such that tonotopic differences for these features are essentially indistinguishable by P14 (latency: base, 5.5±0.2 ms, n=12; apex, 5.6±0.2 ms, n=15 and accommodation (APmax: base, 1±0; apex, 1.1±0.07). However, tonotopic differences in threshold persist (base, -42.3±1.2 mV; apex, -46.7±0.9 mV; P<0.05). Finally, NT-3 application to base neuron cultures promotes lowering of voltage threshold, prolonged latencies and slow accommodation in P6 and P8 neurons. The effect of NT-3 on younger (P3) and older (P10 and P14) neurons is currently being investigated.

Conclusion
During the first 2 weeks of postnatal development, SGNs undergo rapid and considerable alterations in their intrinsic excitability, and NT-3 may play a central role in modulating SGN excitability as the animal matures. Funding: NIDCD RO1 DC01856.

MicroRNA-96 is Required for the Postnatal Refinement of Auditory Nerve and the Survival of Type I Spiral Ganglion Neurons
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Background
Mutations in the seed region of miR-96 cause progressive hearing loss in two DFNA50 families and Diminuendo (Dmdo) mice. Homozygous mutant (Dmdo/Dmdo) mice show profound hearing loss and have extensive sensory hair cell loss at one month of age (Lewis, et al. 2009). Using this animal model, Kuhn et al (2011) demonstrated that miR-96 is essential for morphological and physiological maturation of the sensory hair cells. Although it has been established that miR-96 is
expressed in both sensory hair cells and spiral ganglia (Sacheli et al., 2009; Weston et al., 2011), the pathological alterations of the spiral ganglion neurons (SGNs) and their processes remain to be determined in Dmdo mice.

Methods
Wild type, Dmdo/+ and Dmdo/Dmdo mice were examined at postnatal day (P) 3, P8, P14, P21, P23 and 1, 2 and 3 months of age. SGNs and their peripheral projections were examined using frozen sections and auditory nerve surface preparations. A battery of neuronal cell markers were employed to identify two populations (types I and II) of SGNs and their projections. Synaptic ribbons and the efferent component of the auditory nerve were labeled with antibodies for C-terminal binding protein (CtBP2) and choline acetyltransferase (ChAT), respectively. In addition, ultrastructural characteristics of neurons and fibers in the spiral ganglia of the Dmdo mice were examined by transmission electron microscopy.

Results
Disorganization and increased length (or density) of neurofilament 200 and ChAT positive fibers were seen under inner hair cells of both Dmdo/+ and Dmdo/Dmdo mice aged P14 to P23. Ultrastructural observations indicated several signs of SGN degeneration in Dmdo/+ and Dmdo/Dmdo mice including thinner myelin, loose myelin lamellae, vacuole-like inclusions in the cytoplasm of Schwann cells and “bracelet-like” processes around neuronal cell bodies. Quantitative analysis of cell density in two populations of SGNs shows a significant loss of type I SGNs in the basal turn of the one-month-old Dmdo/Dmdo mice.

Conclusion
MiR-96 is required for the postnatal refinement of the auditory nerve and the survival of type I SGNs. An early onset of type I SGN loss suggests that neuronal degeneration is one of the primary changes contributing to hearing impairment in Dmdo mice. This study was supported by NIH R01 DC7506 (H.L.), NIH P50 DC0422 (H.L.) and NOHR 206048 (D.M.F.).
A Viscoelastic Model of Adaptation in Mammalian Auditory-Nerve Fibers
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Background
Responses of mammalian auditory-nerve fibers to acoustic stimuli show pronounced adaptation and recovery from adaptation. During a tone of constant amplitude, the spike rate declines markedly, particularly during the first tens of milliseconds following stimulus onset. Upon cessation of the tone, the spike rate drops to values below the spontaneous rate and gradually recovers. A variety of functions are ascribed to adaptation, including shifting the cell’s sensitivity range. Most models of the adaptation behavior of mammalian auditory-nerve fibers assume the depletion of pools of synaptic vesicles in inner hair cells. However, recent research in mammalian systems suggests the presence of two alternative adaptation mechanisms, reflected in transducer currents and membrane potentials. A fast mechanism is thought to close transducer channels upon Ca2+ binding, whereas a slow mechanism is thought to relieve tension on the channel through slippage of a myosin motor, both often modeled as exponential decays. Here, we examine whether a double exponential decay model with viscoelastic mechanics, inspired by the work of Howard and Hudspeth (PNAS 1987), can describe the responses of mammalian auditory-nerve fibers over the entire peristimulus time course.

Methods
Spike times were obtained from extracellular recordings of auditory-nerve fibers in barbiturate-anesthetized cats. Stimuli were 100-ms tones of different sound pressure levels at each frequency (mainly characteristic frequency), repeated at least 100 times at a rate of 4 Hz. Prior attempts to characterize adaptation have mainly involved fitting parametric models to poststimulus time histograms (PSTHs). However, such fits produce parameter estimates that depend on the chosen bin width. We instead fit our model to the cumulative spike count to avoid this problem. Cumulative spike count functions maintain maximal temporal precision and thus retain more information than most PSTHs, yet are smooth and allow precise fits.

Results
The viscoelastic model proposed here incorporates fast and slow exponential decay and recovery components, with a comparable number of parameters to classical models. We found that the viscoelastic mechanical model of adaptation can describe the data with small and unsystematic residual errors, throughout the entire peristimulus time interval. Furthermore, the parameters have simple physiological interpretations and seem to show meaningful relationships to stimulus parameters.

Conclusion
The viscoelastic description of adaptation may constitute a viable alternative to the assumption that vesicle depletion and re-supply are the prime factors underlying adaption in mammalian auditory-nerve fibers.

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Assessment of Neural Phase-Locking at the Round Window in Human
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Background
Phase-locking is a fundamental property of the peripheral auditory system. In single auditory nerve fibers, phase locking declines with frequency and becomes undetectable at an upper frequency limit which differs between species. This limit is unknown in humans, and widely differing values are assumed in the interpretation of psychophysical results. Our previous work in cat (Verschooten et al 2011) suggested that mass-potentials at the cochlear round window allow an assessment of the limit of neural phase-locking. Here, we further test this assumption, and examine whether recordings through the middle ear can also be used in macaque monkey and human volunteers to assess phase-locking.

Methods
A minimally invasive transtympanic protocol was developed to record mass-potentials from the cochlear promontory or from the niche of the round window in macaque and human. This involved a custom made ear mold with openings for the transtympanic needle electrode and for acoustic stimulation (ER-2 earphone, calibrated in-situ with an ER-7 microphone). We used a forward masking paradigm to disambiguate contributions of receptor and neural origin: probe tones with alternating polarity were preceded by a forward masker of the same frequency to temporally mask the neural contribution.
in the response. The phase-locked neural contribution was estimated from the difference between the unmasked and the masked responses; the compound action potential was excluded by either filtering or by subtracting the results to stimuli of opposite polarity. To validate the applied method, we compared the results measured at the round window before and after administration of tetrodotoxin (10mM, round window) in cat.

**Results**
An adaptive, neural contribution could be measured on the cochlear promontory of macaque and human. Its magnitude and signal to noise ratio were much lower than in cat (Verschooten et al 2011). A neural, stimulus related phase-locked component could be detected for probe frequencies below 2 kHz in macaque and human. In human, spectral analysis sometimes revealed non-stimulus oscillatory components for probe frequencies above ~1 kHz. In cat, application of TTX showed that the adaptive part of the cochlear mass-potential was reduced by >90% over a wide frequency range, consistent with a neural origin.

**Conclusion**
Neural, phase-locked mass-potentials are recordable with a minimally invasive technique from awake, normal hearing subjects. We found evidence for stimulus-frequency phase-locking below 2 kHz in macaque and human. Supported by the Fund for Scientific Research – Flanders.

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**Examining the effects of vowel training on neural response sensitivity in ferret auditory cortex**

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**Background**

Neurons in ferret auditory cortex are modulated by the pitch, spatial location and spectral timbre of artificial vowels and found to be unable to represent either pitch, timbre or spatial location in a manner that was independent of changes in the other two dimensions. This study compared the responses of animals trained to discriminate artificial vowel sounds with our previous control data.

**Methods**

Three ferrets were trained to identify two of the vowel sounds (/u/ and /ε/) used in the previous study in a two-alternative forced-choice paradigm. Animals were anesthetised and recordings were made from neurons located across five cortical fields. Our stimuli were all possible combinations of four vowel timbres (including the two trained vowels), four fundamental frequencies and four spatial locations (Bizley et al., 2009), as well as pure tone stimuli at varying sound intensities and noise bursts.

**Results**

We quantified to what extent neurons were sensitive to pitch, timbre or location with a variance decomposition approach and compared the neural responses from the trained animals with the control data we have previously published. As in our control data the majority of neurons were sensitive to more than one sound feature. Neurons in the trained dataset were significantly more informative about spatial location of the artificial vowels than responses from control animals and, over all, relatively less informative about the pitch and timbre of the sound. When we considered the responses recorded in each of the cortical fields individually, we observed that timbre sensitivity was significantly lowered in the primary auditory cortex (fields A1 and AAF), but was markedly increased in field PPF for trained animals versus controls.

**Conclusion**

These findings suggest that training elicits field-specific changes in the representation of vowel identity.

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**Determining the Relative Importance of Cues For Concurrent Vowel Identification**

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**Background**

Listeners with normal hearing have the ability to understand one talker in the presence of other talkers. Difference in fundamental frequency (F0) is one important cue for talker segregation. When listening to two vowels simultaneously, identification of both vowels increases as the difference in F0 of the two vowels increases. Although vowel segregation
clearly benefits from F0 differences, listeners’ ability to segregate two vowels with identical F0 suggests that other cues are contributing, such as vowel spectral characteristics. To assess the relative importance and interaction of these and other cues, we measured concurrent vowel identification under conditions that limited the availability of certain cues. A computational model of responses of auditory-nerve fibers was used to predict the neural basis of these cues.

Methods
Younger adults with normal hearing listened to two synthetic concurrent vowels with and without F0 difference at vowel levels ranging from 25-85 dB SPL. Vowel pairs were selected from five vowels and included same and different pairs. In the computational model, F0 coding using a periodicity metric across various fibers, and formant coding using a synchronized rate metric, were predicted for vowel pairs at each level. These predictions assessed the extent to which (1) reduced salience of F0 difference cues at lower levels can be attributed to poorer phase locking of auditory-nerve fibers to harmonics of the two F0s and (2) changes in vowel identification with increasing level in the absence of F0 difference cues can be attributed to changes in phase locking of auditory-nerve fibers to vowel formants.

Results
Identification of both vowels improved as vowel level increased from low- to mid-levels and then declined at higher levels. The slope of the functions relating vowel identification to vowel level was differentially affected by vowel type and F0 difference. At most levels, identification of both vowels was better with than without an F0 difference. In contrast, scores for vowels with and without F0 difference were more similar at lower levels, suggesting a reduced contribution of this cue.

Conclusion
Younger adults with normal hearing use F0 difference, spectral, and other cues to identify two concurrent vowels. The relative importance of these cues for each vowel varies with vowel level and may relate to level-dependent changes in phase-locking of auditory-nerve fibers to vowel harmonics and formants.

[Work supported by NIH].

The contribution of low spontaneous rate, auditory nerve fibers to recognizing speech in noise

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Background
Humans are remarkably good at resolving speech in noise. While humans still perform well at -12 dB signal-to-noise (SNR) ratios and lower, even the best computer automatic speech recognition systems have significant performance decrements at around 0 dB SNR. How humans achieve this robust performance remains unclear. There is evidence that auditory nerve fibers with low spontaneous rates, which have been traditionally thought to be most important for encoding a large dynamic range, may play an important role (e.g. Silkes and Geisler, 1992). This proposal has been explored primarily for frequency cues, but not for the temporal cues that have been described recently (e.g. Clarey et al., 2004).

Methods
In this study, we presented speech consonants in quiet and with background noise to ketamine-anesthetized chinchillas while recording from individual auditory nerve fibers. A peristimulus time histogram (PSTH) was computed from the responses of each auditory nerve fiber to each consonant, and these PSTHs were averaged across all fibers to create the consonant’s ensemble response.

Results
While this method produced an ensemble response which displayed a unique temporal pattern for each consonant in quiet, these temporal patterns were not apparent for consonants played in noise. Using only the responses from low spontaneous rate fibers, however, allowed for a reasonable recovery of these temporal patterns.

Conclusion
It appears that low spontaneous rate fibers are important for carrying the temporal cues when a consonant is heard in noise and would be crucial for robust speech recognition in noise.

This research was supported by the National Science Foundation.
Voltage-gated ionic conductances required for action potential firing in auditory nerve fibers
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Background
Depolarization of the inner hair cell triggers glutamate release onto the dendrite-like processes of spiral ganglion neurons (SGNs) and drives action potentials, which are convey to the brain. Whereas knowledge of the transfer function at the ribbon synapse has considerably progressed, little is known about the voltage-gated ionic channels which shape the action potential firing. Here, we provide a comprehensive computational model bridging the gap between the voltage-dependent currents measured in vitro on fresh isolated SGNs and spikes (extracellular action potentials) recorded in vivo from guinea pig auditory nerve fibers.

Methods
Voltage-dependent currents (Na+ and K+) of SGNs somata patch-clamp recordings were fitted by a Hodgkin-Huxley model with a full-trace fitting Williams algorithm. Node of Ranvier model was design from the hypothesis that channel expressed on soma were identical, but differ in density. Simulated spikes were adjusted in order to match in vivo single-unit recordings with gradient-descent algorithm.

Results
Computation of the data allows to the identification of: i) one fast inward Na+ current (GNa activation: V1/2=-33 mV, &>=0.5ms; inactivation: V1/2=-61 mV, &=< 2ms); and ii) two K+ delayed-rectifier conductances, a low voltage-activated component (GKL, activation: V1/2=-56 mV; &=< 5 ms) and a high voltage-activated component (GKH, activation: V1/2=-41 mV; &>=2.5 ms). Node of Ranvier model generate spikes that fit with in vivo recordings. Interestingly, the different spike durations along the tonotopic axis measured in vivo (i.e. 450 ms peak-to-peak duration versus 250 ms for 1 to 20 kHz, respectively) was explain by a gradual change in Na and K channel densities along the cochlea (GNa ~78 nS, GKL ~9 nS, GKH ~3 nS at 1 kHz versus GNa ~90 nS, GKL ~12 nS, GKH ~6 nS at 20 kHz).

Conclusion
This study identifies the ionic conductances and densities, which shape the action potential waveform of auditory nerve fibers and suggests that the interplay of fast inward Na+ current and the two K+ delayed-rectifier enables the auditory nerve fibers to sustain high firing rates.

Merlin is expressed in spiral ganglion neurons (SGNs) and inhibits neurite growth
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Background
Merlin, the product of the nf2 tumor suppression gene, mediates cell-cell contact information to inhibit cell proliferation and motility. It does so by inhibiting several progrowth intracellular signaling cascades such as Ras, Rac, phosphatidylinositol-3 kinase/Akt, extracellular regulated kinase, and c-Jun N-terminal kinase, which have also been implicated in neuritogenesis and neurite growth. We hypothesize that neuronally-expressed merlin regulates neurite growth by inhibiting these signals in response to cell-cell contact cues.

Methods
Immunostaining of cochlear organotypic explants with anti-merlin antibodies was used to verify merlin expression in SGNs and their peripheral neurites. We used a tamoxifen (Tx)-inducible cre-mediated recombination system to knock out merlin in neonatal SGN, trigeminal ganglion (TGN), and cortical neuron cultures. Dissociated cultures were maintained in the presence or absence of Tx for 24h to induce Cre expression. Neurite length was measured after an additional 48h in culture. To distinguish between the effect of loss of merlin expression in SGNs and loss in Schwann cell (SCs), we plated neurons that had been treated or not treated with Tx onto a bed of SCs that had likewise been treated or not treated with Tx.

Results
We find that merlin is expressed in SGNs raising the possibility that merlin regulates SGN neurite growth in addition to its well-known role in SC tumorigenesis and cell motility. Reduction of merlin expression in neurons but not SCs increases both the number and length of neurites in SGNs and TGNs. Decreasing merlin expression via simultaneous treatment of SCs and neurons with tamoxifen increased neurite length from 365+/−125µm (mean +/- SE) to 522+/−165µm in cultured
neonatal SGNs and from 372+/−31µm to 510+/−113µm in TGNs. SGN and TGN neurite length in samples where SCs were treated with tamoxifen but not neurons was 388+/−122µm and 391+/−101µm respectively. SGN and TGN neurite length in samples where neurons were treated with tamoxifen but not SCs was 500+/−92µm and 511+/−93µm respectively.

**Conclusion**

Inhibition of neurite growth by merlin may be due to the effect of merlin expression in neurons rather than in SCs. Current studies examine the merlin-sensitive signaling pathways that contribute to SGN neurite growth and the influence of merlin status on synaptogenesis. This raises the possibility that merlin regulates neural development and/or regeneration, in addition to its role in suppressing tumor growth.

**The Organ of Corti’s Role in Regulating Spiral Ganglion Neuron Axonal Projection Length**

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**Background**

The distinct morphological features of a neuron can affect its signal transmission. Spiral ganglion neurons (SGNs) display a bipolar phenotype in which specific features are graded, such as soma area, axon diameter, and branching patterns. For example, the axon segment from the foramen nervosum to the Schwann glial border SGB (intracochlear axon) is longer for low frequency neurons (apex) relative to high frequency neurons (base) whereas, the segment from the SGB to the bifurcation (root branch) shows the opposite pattern. The length differences between these two regions compensate for each other to the extent that the total length from the organ of Corti (OC) to the bifurcation is the same for apical and basal neurons (Fekete D.M. et al., JCompNeuro, 1984). This study examines the axon length that extends from the soma to the region of axon bifurcation when co-cultured with regions of the OC.

**Methods**

The preparation pairs the hair cells and their surrounding satellite cells (micro-isolates) with SGN explants (Flores-Otero et al., JNeurosci, 2007). Immunolabeling with α-β-tubulin antibody was utilized to identify the neurons; axon length was measured in IPlab (Scanalytics) from the soma to the first branch point.

**Results**

Here we examine the role that OC micro-isolates play in regulating axon length. Our in vitro data recapitulated the axon length relationship of the intracochlear axon measured in vivo. When apical hair cell micro-isolates were paired with apical neurons the measured axon length was significantly longer (779 ± 70µm, n=17) when compared to basal hair cell micro-isolates paired with basal neurons (462 ± 82µm, n=10; p<0.01). The peripheral targets were identified to regulate this parameter when the OC micro-isolates taken from different tonotopic regions were mixed and matched with SGNs. The apical hair cell micro-isolates co-cultured with basal neurons increased axon length (903 ± 75µm, n=7; p<0.01) when compared to basal hair cell micro-isolates co-cultured with basal neurons. Conversely, basal hair cell micro-isolates co-cultured with apical neurons reduced axon length (398 ± 60µm, n=3; p<0.05) relative to apical hair cell micro-isolates co-cultured with apical neurons.

**Conclusion**

The intracochlear axon length relationship is reflected in our measurements. Thus, elements from the OC can regulate this parameter suggesting that the intracochlear axon and the root branch length are differentially regulated. NIH NIDCD R01 DC-0185.

**Improved Parameters and Expanded Simulation Options for a Model of the Auditory Periphery**

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**Background**

The auditory periphery impacts the neural representation of speech and is the sole conduit for information to reach the higher auditory centers. A phenomenological model of auditory-nerve (AN) responses was developed to understand the staged transformations in the auditory periphery (Zilany et al., 2009, JASA, 126:2390). This model incorporates most of the nonlinearities observed in AN responses; however, it does not accurately simulate data from Liberman (1978, JASA 63:442) describing the discharge rate at saturation as a function of characteristic frequency (CF) for higher CFs. The responses of higher-CF model fibers to low-frequency tones (e.g. 500 Hz) are also erroneously much higher than the
responses of low-CF model fibers (Fig. 1A). This issue was addressed here, and the model was also extended to include a set of parameters for a human AN model.

Methods
It was determined that the source of the mismatch between the reported saturation-rate data from Liberman (1978) and the model responses lies in the parameters for the model stage that describes synaptic adaptation. With minimal effects to other response characteristics, the parameters of the synapse model were modified to solve the problem described above. Additionally, an analytical method was adopted from Vannucci and Teich (1978, *Optics Communications* 25:267) to compute the instantaneous mean discharge rate and variance from the model’s synapse output that takes into account the effects of refractoriness. Finally, a set of parameters that describes estimates of human tuning is now included (Ibrahim and Bruce, 2010, *The Neurophysiological Bases of Auditory Perception*, pp. 429) in addition to parameters for cat.

Results
The new model with the modified set of parameters has been extensively tested against the published and recorded data from the AN. The saturation rates, especially for high-CF model fibers, were corrected by the modification (Fig. 1B), whereas other response properties were largely unchanged. The forward-masking properties were affected to a small degree, as could be expected after modifying synaptic adaptation.

Conclusion
The revised AN model is a better candidate to examine realistic neural-encoding hypotheses, especially those involving high CFs. The changes made to the previous version correct the saturation rates as a function of CF without adversely affecting other response properties. The revised model also includes a set of parameters that describes the middle ear transfer functions and peripheral tuning in human.

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FIG 1. Response area to a 500 Hz tone (50 ms duration with 3.5 ms on-off ramp). Mean rate responses of the 109 AN fibers with CFs logarithmically spaced from 500 Hz to 20 kHz (along x-axis) are plotted for sound levels ranging from -15 to 120 dB SPL (along y-axis) in steps of 2.5 dB. A) Responses from the Zilany et al. (2009). B) Responses from the new model.
Amplitude of masked compound action potentials indicates location of laser-induced lesions in the auditory periphery
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Background
Auditory nerve degeneration is thought to lead to difficulties with understanding speech, especially in difficult listening situations. Recent research suggests that the presence of neural degeneration may be missed if threshold and latency are the only metrics used to assess auditory evoked potentials. Adding amplitude as a metric, overall nerve survival can be inferred from high-level compound action potentials (CAPs). However, the inherent variability in absolute amplitude measures may confound the prediction of the extent of nerve damage. Our recent experiments have focused on circumventing this confound and improving the location-specificity of damage predictions by tracking CAP amplitude growth while systematically limiting the region of synchronously firing auditory neurons with a high-pass masking paradigm. The objective of this study was to determine if the pattern of CAP amplitude growth in animals with peripheral auditory nerve lesions indicates the corresponding cochlear location of the lesion.

Methods
Mongolian gerbils with laser-induced lesions in the auditory periphery were compared to normal-hearing gerbils. CAPs were evoked with broadband chirps at 90 dBSPL during simultaneous masking noise high-passed in 1/3 octave intervals between 0.4 and 50 kHz. Cumulative amplitude functions were constructed by plotting normalized CAP amplitudes as a function of distance from cochlear apex (inferred from masker cutoff frequencies). Animals were euthanized and the cochleae were fixed, decalcified, and sectioned into three turns. The cochlear turns were first stained with phalloidin to enable imaging of hair cells under confocal microscopy. The peripheral processes of the auditory nerve were then stained with osmium and imaged under light microscopy.

Results
Confocal images of laser-exposed cochlea revealed circular lesions through regions of the osseous spiral lamina and missing hair cells. Light micrographs of osmium-stained cochleae revealed regions of severed neurons and lighter staining consistent with myelin damage. Cumulative amplitude functions in normal-hearing gerbils were sigmoidal in shape and approached saturation at the basal end of the cochlea. In lesioned gerbils, the amplitude functions plateaued in regions that generally corresponded to the cochlear region at which the laser was directed.

Conclusion
These data indicate that the pattern of CAP amplitude growth may identify the cochlear location of auditory nerve damage. Future experiments gauging the reliability of amplitude growth functions and enhanced anatomical measures (e.g., dual-fluorescence labeling of hair cells and neurons) are needed to determine the predictive power of this technique.

Supported by Action on Hearing Loss, UK

Heterogeneous Electrophysiological Properties from Murine Spiral Ganglion Neurons in Acutely Prepared Cochlear Slices
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Background
Type I spiral ganglion neurons (SGNs), composing most of the ganglion (90-95%), are responsible for conveying the acoustic signals that are encoded by the inner hair cell (IHC) receptors to the central auditory pathways in the form of action potentials. Although the initiation of action potential occurs at the synaptic terminal that innervates the IHC, the cell soma from acutely dissociated neurons is excitable and thus capable of shaping the signal during its propagation. Moreover, the timing and intrinsic excitability features of cultured SGNs from one week old animals are distinctively organized along the tonotopic axis and may relate to coding separate sub-modalities of acoustic features (Liu and Davis, J Neurophysiol, 2007). To extend the studies in culture and facilitate our understanding of spiral ganglion function ex vivo, we have begun to characterize the electrophysiological properties from neurons in acutely prepared cochlear slices and have found evidence of the heterogeneity and firing features that are consistent with observations made in vitro.

Methods
After removing the cochlea, slices (Jagger et al., J Neurosci Methods, 2000) were prepared on a vibratome (Leica VT1200, ~200 µm slice thickness) from postnatal day 4 (P4) to 8 (P8) CBA/CaJ mice. SGNs were visualized under...
infrared differential interference contrast (IR-DIC) microscopy (Olympus BX50WI) and recorded with the whole cell patch clamp technique.

Results
The SGNs from slices fire well formed action potentials (amplitude: 87 mV ± 4 mV, n = 8). Both rapidly and slowly accommodating neurons were observed during a 240 ms step current stimulation: two cells fired a maximum of 13 and 16 action potentials, respectively, and the rest fired fewer than 7 (2.2 ± 1.2, n = 5). A heterogeneous distribution was found for action potential latency at threshold (ranging from 4.1 to 9.9 ms; 7.8 ± 1.4 ms, n = 8) and voltage threshold (-60.01 to -48.19 mV; -55.1 ± 1.4 mV, n = 8); all features that are consistent with our culture recordings.

Conclusion
The basic firing features including accommodation, action potential latency at threshold and voltage threshold from spiral ganglion neurons in acutely prepared slices are heterogeneously distributed and similar to postnatal SGN recordings in culture. Future experiments will address whether the observed heterogeneity reflects tonotopic changes or developmental influences from P4 to P8. Supported by NIH NIDCD RO1-DC01856.

Noise-induced hearing loss increases sensitivity to fast temporal modulations in the auditory nerve: evidence from Wiener kernel analysis
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Background
The increase in the bandwidth of cochlear filtering associated with many forms of hearing loss should theoretically extend temporal modulation sensitivity to higher modulation frequencies. However, both behavioral studies in humans and neurophysiological studies in animals have found little evidence of enhanced sensitivity to fast temporal modulations.

Methods
Here in anesthetized chinchillas, we estimated temporal modulation transfer functions (TMTFs) from auditory-nerve fiber responses to near-threshold Gaussian noise stimuli. TMTFs were computed from second-order Wiener kernels of the neural responses, and compared between control animals (N=12) and animals with noise-induced hearing impairment (N=16) to assess possible changes in sensitivity to fast temporal modulations. We studied temporal modulation tuning in 163 fibers with characteristic frequencies (CFs) between 0.25 and 10 kHz.

Results
The noise exposure caused a loss of sensitivity and increase in the bandwidth of the pure-tone tuning curve across the full range of CFs investigated. The corner frequency of the TMTF measured -3 dB and -6 dB from the peak of the function increased with CF in both the noise-exposed and control groups, as observed previously in studies using neural responses to sinusoïdally amplitude modulated (SAM) tones. In contrast to previous studies however, the corner frequency of the TMTF was 20-30% higher in noise-exposed fibers than in control fibers at any given CF. Previous studies with SAM tones showed little effect of noise-induced hearing loss on TMTF corner frequency. The increase in TMTF corner frequency observed here was generally less than the increase in the bandwidth of the pure-tone tuning curve, which increased by 100-200% relative to controls in some cases.

Conclusion
The results suggest that noise-induced hearing loss does in fact increase neural sensitivity to fast temporal modulations in the auditory periphery. The discrepancy with previous neurophysiological results probably reflects differences in the choice of stimulus. Whereas previous work relied primarily on SAM tones, we estimated sensitivity to temporal modulation using a dynamic stimulus with broadband acoustic structure similar to many natural sounds including speech. Inasmuch as the results demonstrate a deviation from normal spatiotemporal neural coding of sound in the cochlea, they may help explain the basis of degraded speech perception in people with common forms of hearing loss. Furthermore, the results underscore the value of studying hearing loss with more natural, real-world acoustic signals.

Research supported by NIH (NIDCD) F32-DC012236 and R01-DC009838.
Correlation and Temporal Properties of the Auditory Nerve
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Background
Modeling of the spontaneous activity in auditory nerve fibers failed if an exponential (Poisson) or gamma distribution of the inter-spike time-intervals was assumed. Nevertheless, the distribution may still be described by a stochastic process. Spontaneous and driven activity of pigeon auditory nerve fibers was tested for a renewal process. The experiments falsified mutually statistically independent lengths of time intervals between successive action potentials. The question for this study was if the generation of action potentials on the auditory nerve may be described by a doubly embedded stochastic model, the Hidden Markov Model (HMM), and whether the model parameters change for various inputs at different levels.

Methods
The analysis is an extension of the work by Richter et al., 1996. Spontaneous activity and responses to sinusoidal and white noise acoustic stimuli, which were recorded from single pigeon auditory nerve fibers, were used to estimate the transition and emission matrices of the HMM. Newly generated sequences of events (surrogate data) were generated and the autocorrelation functions of the original data and the surrogate data were compared.

Results
Autocorrelation functions of the original data and surrogate data were not significantly different. Although action potential repetition rate was below 100 Hz, fine structure of more than 1 kHz could be extracted from the spike trains. This fine structure was maintained in the surrogate data despite the fact that little correlation existed between the occurrence of the action potentials (recorded data) and the surrogate data. Fine structure of responses could be modeled with low event repetition rates.

Conclusion
The HMM provides a method to create sequences of events that contain similar high frequency structure as the original sequence of action potentials. The HMM might be used as a spike generator for auditory prostheses to stimulate the auditory system at low pulse repetition rates maintaining high frequency fine structure. This is particularly attractive for infrared neural stimulation.

Unstable representation of sound: A biological marker of dyslexia
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Background
Learning to read proceeds smoothly for most children, yet others struggle to translate verbal language into its written form. Poor readers often have a host of auditory, linguistic, and attention deficits, including abnormal neural representation of speech and inconsistent performance on psychoacoustic tasks. We hypothesize that this constellation of reading-related deficits arises from the human auditory system failing to respond to sound in a consistent manner, and that this inconsistency impinges upon the ability to relate phonology and orthography during reading.

Methods
102 normal hearing children (ages 6-13) were classified as good (> 120), average (105-95), and poor (< 90) readers based on their scores on a test of reading fluency (Torgesen et al., 1999). We collected evoked auditory brainstem responses to [ba] and [ga] stimuli and calculated response consistency by correlating responses from the first 3000 repetitions of the stimuli to those from the last 3000 repetitions, with a higher r-value representing more consistency (conversely, less variability).

Results
Poor readers have significantly more variable auditory brainstem responses to speech than do good readers. The groups do not differ on prestimulus response amplitude, indicating differences in response consistency were independent of physiological noise levels.

Conclusion
Here we provide evidence that poor readers have less stable auditory nervous system function than do good readers. Although causality cannot be determined, heightened variability in nervous system function may underlie fluctuations in
directed attention and impaired speech understanding due to inconsistent encoding. Our results suggest that good readers profit from a stable neural representation of sound, and that children who have inconsistent neural responses are at a disadvantage when learning to read.

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Multichannel ABR.
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Background
The auditory brainstem response (ABR) is widely used clinically in humans, and in animal models, to assess the integrity of the peripheral auditory system. The signals are extremely small, embedded in environmental and physiological noise of much higher amplitude, and measurements require many repetitions and are time-consuming. Our hypothesis is that the method can be improved by increasing the number of recording electrodes, and processing the data with recent multichannel signal processing techniques.

Methods
Data are acquired using a standard 32-channel EEG system (Biosemi). Multiple electrode layouts are investigated in an attempt to (a) optimize sensitivity to brainstem components, and (b) optimize coverage of major noise sources, so that they can be factored out. Data are processed using the Denoising Source Separation (DSS) algorithm, that finds the optimal combination of channels to maximize repeatable activity. Data have also been acquired in mouse using multiple electrodes.

Results
Compared to standard ABR (two or three electrodes), multichannel ABR produces data with a better signal-to-noise ratio (SNR). In many instances multiple components may be observed, with distinct time-course and spatial signature, suggesting that it may be possible to differentiate contributions from different processing stages within the brainstem.

Conclusion
Multichannel recording and processing techniques improve the signal-to-noise ratio and resolution power of ABR measurements.

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Subcortical Pitch Encoding of Speech Sounds in the Normal and Impaired Auditory Systems
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Background
Several perceptual studies have demonstrated that hearing impaired (HI) listeners have reduced access to temporal fine structure (TFS) information, which may possibly be due to reduced phase-locking ability. However, there are few electrophysiological studies examining phase-locking in HI listeners, particularly at the subcortical level using the frequency following response (FFR). The FFR is a scalp recorded sustained potential, reflecting phase-locked activity from a population of neural elements in the rostral brainstem. Previous FFR studies have found that the phase-locking ability to second formant transition in HI individuals is significantly reduced as compared to normal hearing (NH) listeners. Here we examine brainstem neural phase-locking (as measured by the FFR) to a synthetic steady state vowel in NH and HI (mild-moderate SNHL) listeners. First, we aim to characterize phase-locking to the envelope and TFS in both groups. Secondly, to determine if the difference between the two groups is related to audibility, we measured FFRs at multiple sound pressure levels (SPLs) to facilitate comparisons between the two groups at equal sensation levels (SLs) and equal SPLs.

Methods
FFRs were recorded from NH and HI listeners in response to a steady state synthetic vowel at multiple sound pressure levels. Temporal and spectral analyses were conducted to describe the FFRs. Pure tone audiograms, speech-in-noise testing and case-history information was obtained for all subjects.

Results
Preliminary results indicate statistically significant differences between the NH and HI groups for phase-locking to both envelope and TFS. Comparisons at equal SLs and SPLs continue to show differences between the two groups. Also
notable is the variability noted within the HI group: a combination of clinical testing, demographic and case-history information collected seems to suggest a variety of factors (age, pure tone thresholds, use of amplification, etc.) could be contributing to the intra-group variability.

Conclusion
FFRs in NH are more robust in amplitude and phase-locking than those in HI listeners. Audibility does not appear to be the chief cause underlying the differences seen between the two groups. These results are consistent with those obtained in previous work by Plyler and Krishnan (2001). A multitude of other factors may be causing the intra group variability seen in the HI group. The results observed in the hearing impaired FFRs may be due to disruption in temporal synchronization of neural activity across a population of neural elements.

Context-dependent Processing of Shepard Tones in the Human Auditory Brainstem and Cortex - an Electrophysiological Study
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Background
Recently, a series of studies using the frequency-following auditory brainstem response (FFR) have shown that the encoding in the human inferior colliculi is not merely determined by the acoustics of the incoming stimulus, but can be enhanced by short-term experience with specific auditory material, e.g. stimulus repetition. Here, we tested whether stimulus regularity, as in the case of regularly presented Shepard tones leading to the illusion of a continuously descending pitch, will lead to a modulation of the frequency representations in the brainstem.

Methods
23 healthy human subjects listened to sequences of tones presented monaurally through headphones to the right ear (75 dB SPL; SOA of 240 ms) while the left ear was masked with white noise. Sequences consisted of Shepard tones that were presented continuously either in a descending or in a random manner, the later with the restriction of keeping the step-size from one tone to the next identical to the one in the descending condition. EEG was recorded from Cz with an electrode at the left earlobe serving as reference, sampled at 20 kHz, and analyzed for the auditory brainstem response (ABR). Additionally, middle-latency responses (MLR) and long-latency responses (LLR) of the auditory evoked potential (AEP) were obtained to study later regularity-related effects.

Results
In the ABR time range, FFR to the prominent partials of the Shepard tones were smaller in amplitude in the descending than in the random condition (p=0.029). Results did not reveal an interaction between conditions and the amplitudes with which single partials were represented in the brainstem FFR. In the MLR time range, tones in the random condition elicited a smaller Pa-Nb complex than tones in the descending condition (p=0.030). LLR potentials in the time range from 130-180 ms were more negative in the random than in the descending condition (p=0.001).

Conclusion
In the long-latency range, results resemble a typical surprise-related negativity for randomly presented tones. In contrast, in the earlier latencies of the AEP, the pattern for regular descending Shepard tones (attenuated brainstem FFR, enhanced MLR) is opposite to the pattern observed for repetition regularities in previous studies (enhanced FFR, attenuated MLR). This could suggest that earlier processing stages are rather influenced by the higher local probability of tone repetitions in the random condition of the present study and not by the non-repetitive stimulus regularity.

Rapid Acquisition of Auditory Brainstem Frequency Following Responses
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Background
Auditory Brainstem Frequency Following Responses (FFRs) provide a non-invasive measure of encoding in the auditory pathway up to the Inferior Colliculus and have been used extensively both clinically and in basic neurophysiological investigation of auditory function. FFR data acquisition for simple stimuli such as click trains, tones, and speech tokens typically involves thousands of presentations of each stimulus type, sometimes in two polarities leading to acquisition times exceeding an hour per subject. Since the FFR sources are deep compared to the primary noise sources (i.e.)
cortical activity, FFR varies more smoothly over the scalp than cortical noise. Additionally, since the FFR is a multi-source mixture, there are phase differences in the responses at different scalp locations. We present a novel approach to exploit the above properties using multichannel recordings and an eigendecomposition of the complex cross channel spectral density (CSD) matrix (i.e.) complex Principal Component Analysis or cPCA to extract FFRs efficiently.

Methods
First, we use simulations to validate the cPCA method. 32 channels, each containing a 200 ms burst of a 100 Hz sinusoid with a random phase, were added to 400 epochs of background EEG data to mimic SNR and between channel noise covariance of a real multichannel FFR recording. A multi-tapered estimate of the 32x32 normalized CSD matrix C(f) (complex coherence matrix) was diagonalized using eigendecomposition. The function d(f) of the highest eigenvalue in each frequency bin provides the inter-trial phase coherence of the first principal component (the extracted FFR) with the corresponding 1x32 eigenvector v(f) providing the complex weighting function to combine the different channels with appropriate phase lags. An identical analysis is then applied on 32 channel EEG data acquired from human subjects stimulated with 400 trials of 200ms click-train bursts at 70 dBSPL in 2 polarities to extract FFR components phase locked to the envelope in the different cochlear channels.

Results
The cPCA eigenvector accurately estimated the phase lags needed to align the FFR across channels in the simulated data with d(f) showing a very robust 100Hz peak compared to either an individual channel or traditional time-domain PCA. Preliminary results with real data suggest that a fifth of the trials can be combined to yield an SNR comparable to using all the trials with just the best electrode in the phase-locking measure.

Conclusion
The proposed method makes rapid acquisition of FFR data possible.
Hearing Threshold Estimation by Linear Regression on Auditory Brainstem Parameters in Rats
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Background
The hearing threshold is usually estimated by presenting a sequence of decreasing intensity auditory clicks with predetermined step size (4, 5 or 10 dB) until some or all the Auditory Brainstem Response (ABR) waves disappear within the physiological and electrical noise present in the system. This method is usually refined by further exploring the estimated threshold with a smaller intensity step size. These procedures exhibit the inconvenience that the noise level varies from one setup to another and with time.

Methods
20 Sprague Dawley rats weighing 300 ± 50 grams were used according to the local experimental animal research regulatory authority. The ABRs were obtained under xylazine-ketamine anesthesia (6-87 mg/Kg) administered intraperitoneally. The animal body temperature was kept at 37 ± 0.5 C. The stimuli, consisting in compression clicks 100 μS in duration, were presented at a repetition rate of 50 Hz inside of a custom-made sound attenuated booth (50 x 30 x 30cm) exhibiting more than 35 dB attenuation above 1 kHz against the ambient noise. The stimulus sound intensity was varied in a predetermined pseudorandom sequence covering the hearing range of the rat, with two successive steps not differing more than 20 dB. The stimuli were presented binaurally by a commercial speaker placed 17 cm right above the rat's head. A specially designed amplifier was used with 20K overall gain between 100 and 1500 Hz (-3 dB) and a 60 Hz notch filter.

The proposed method for the hearing threshold estimation is based on the linear regression of the stimulus intensity versus a calculated parameter of the ABR that conveys enough information related to the response to sound intensity and that exhibits an approximated linear relation with it. We used the RMS and total difference voltages of the ABR. The hearing threshold can be estimated as the stimulus intensity value at which the derived parameter would be zero as if there was no noise. The two estimation results were combined by the weighted average method.

Results
The hearing threshold values obtained by the proposed method are consistently lower than those obtained by observing the extinction of the ABR waves. The difference between the measured values was 5 ± 2 dB (90% C.L).

Conclusion
The proposed regression method represents an objective alternative, consistent with the conventional methods for hearing level threshold estimation in rats which requires no intervention by the operator or researcher and that can be easily automated.
High-Functioning Autistic Children Have Abnormal Stapedial Reflexes

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Background
Autism is a neurodevelopmental disorder, characterized by social and communicative impairments, sensory abnormalities and restricted, repetitive behaviors. The incidence of autism is approximately 1:88 children and 1:54 in boys. Throughout the autistic brain, there is evidence of developmental dysregulation of neurogenesis, abnormal neuronal maturation, including aberrant neuronal migration, alterations in cell body size, number and dendritic morphology. Additionally, abnormalities in auditory processing are present, in most if not all autistic subjects and abnormal brainstem responses implicate dysfunction of the superior olive (SOC). Finally, we have provided evidence for significant and consistent dysmorphology in the SOC of the autistic brain. Thus, we hypothesize abnormalities in the acoustic stapedial reflex (ASR) in autistic subjects and further, that this objective measure of brainstem function can be used as an early screening tool to identify neonates at risk of carrying a diagnosis of autism.

Methods
The ASR utilizes a brainstem circuitry involving the auditory nerve, the ventral cochlear nucleus, SOC and stapedial motor neurons. Based on the known anatomical and functional auditory deficits in autism, we have compared the ASR in neurotypical and autistic children. All control and autistic subjects in this study had normal hearing sensitivity. Control subjects were only included if: they had no family members diagnosed with autism, normal hearing sensitivity, no upper respiratory disorders affecting middle ear function and satisfactory speech/language development. ASRs were measured ipsilaterally and contralaterally at 500, 1000 and 2000 Hz, using .02mI negative deflection as threshold. ASR thresholds, amplitudes and latencies were compared using JMP10.

Results
Overall, autistic subjects have lower ipsilateral ASR thresholds than neurotypical controls. Contralateral threshold differences are not statistically significant and there are no differences in ASR amplitude between autistic and control subjects. However, we observe significant differences in ASR response latencies. Overall in control subjects, ipsilateral latencies are shorter than contralateral responses, but we observed no such contrast in autistic subjects. Further, we observed significantly longer latencies in autistic subjects and significant asymmetry between responses in the left and right ears.

Conclusion
Our previous reports of dysmorphology in the autistic auditory brainstem along with reported auditory dysfunction and our current report of dysfunction in the ASR provide clear evidence of disruption of auditory pathways in the autistic brain. Based on this study, we believe that non-invasive, objective measures of auditory function have utility as a neonatal screening tool to identify patients with a high risk of autism.

Predicting Hearing Status from Auditory Brainstem Responses Differs in Term and Preterm Infants

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Background
Because all newborns are now screened for hearing loss, diagnostic auditory brainstem response (ABR) testing is performed on more infants than ever before. Broadband and frequency specific stimuli are used to obtain information
about auditory neural integrity and sensitivity, respectively. ABR latencies are compared to normative data from decades-old research comprised primarily of neurologically normal preterm infants or term infants with presumed normal hearing. The population of infants undergoing hearing testing today, especially preterm infants, is vastly different than that tested in previous decades. To better understand infant ABRs, a comprehensive model that includes middle ear measures, otoacoustic emissions and outcome measures is needed. The purpose of this study was to take an initial step towards that goal by modeling retrospectively collected ABR data in preterm and term infants with normal hearing (NH), conductive hearing loss (CHL) or sensorineural hearing loss (SNHL). Two hypotheses were tested: (1) Do ABR Wave V latency and Waves I-V interwave latency for infants with NH, CHL or SNHL have different trends as a function of post-conceptional age (PCA)? (2) Is regression different for preterm and term infants?

**Methods**

Two data sets were combined: cross-sectional data from 84 ears of 71 infants who underwent a clinical research protocol; data from 83 ears of 44 infants with 1-3 clinical ABR tests following newborn hearing screening failure. ABR latencies as a function of PCA were modeled using simple, linear regression. Mean data from Eggermont and Salamy (1988) were used as a comparison.

**Results**

In term infants, the y-intercepts for Wave V latency were different for infants with NH, CHL and SNHL, as expected from previous literature. The slope and y-intercept of Wave V regressions for preterm infants differed from term infants. Waves I-V interwave latencies tended to be longer for preterm infants, which differs from Eggermont and Salamy (1988).

**Conclusion**

The results suggest that ABR latency as a function of post-conceptional age can be modeled using regression for ears with NH, CHL and SNHL. Moreover, regression may be different for preterm and term infants. These results provide a basis for further refinement of a model in which middle ear measures, otoacoustic emissions and outcome measures are included. [Research funded by the March of Dimes Birth Defects Foundation (BP), internal research funds (LH), and Vanderbilt University Summer Research Program.]

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**Envelope coding in humans measured with frequency following responses**

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**Background**

The scalp recorded human frequency following response (FFR) represents sustained phase-locked neural activity among a population of neurons in the rostral brainstem. The present study examined the robustness of envelope coding with increasing sound levels as reflected in the FFR component corresponding to the stimulus envelope.

**Methods**

FFRs were recorded from normal-hearing listeners to sinusoidally amplitude-modulated (SAM) tones, transposed tones, and tone complexes of different sound intensities. The SAM tones were 100% modulated; the transposed tone envelopes were half-wave rectified sinusoids. Both SAM and transposed tones had a modulation frequency (fm) of 200 Hz and a carrier frequency (fc) was of 4 kHz. The tone-complexes had frequency components ranging from 3400 Hz to 4600 Hz with an f0 of 200 Hz. Sound intensity was varied from 50 dB to 90 dB SPL in 10 dB steps. Phase-locking-value (PLV), a measure of the consistency of the evoked signals phase at the envelope frequency, was estimated from the FFR component corresponding to the stimulus envelope.

**Results**

Preliminary results indicate that at moderate sound levels, envelope coding (as reflected in PLV measures) was strongest for transposed tones, second best for tone-complexes, and worst for SAM tones. Envelope coding across all three stimuli first improved with increases in sound level, but dropped from 80 to 90 dB SPL.

**Conclusion**

Previous single-unit neurophysiology studies in animals have reported that the envelope coding at the auditory nerve is strongest near threshold and falls dramatically with increasing sound levels. In contrast, these preliminary results suggest that the envelope coding measured from a population of neurons (as reflected in FFRs) remains fairly stable over a broad range of increasing sound levels.
Crossmodal interactions between audition and nociception
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Background
Research in perception has largely focused on bottom-up processing. Interest in the contribution of efferent pathways is relatively recent and these pathways are poorly understood. Refuting the traditional view that subcortical processing is purely automatic, plasticity has been demonstrated in human auditory brainstem. Such efferent modulation may play a key role in cognitive strategies of pain reduction, such as music-induced analgesia. To date, the underlying mechanism of such modulations is largely unknown.

Methods
We investigated crossmodal interactions between audition and nociception. We anticipated that an auditory task would modulate pain perception, and expected that pain would interfere with auditory processing. Like pain, audition can under certain circumstances serve as a ‘danger detection’ mechanism. We predicted that the earliest component of this crossmodal modulation is mediated by a low-level pathway, subserving a combined auditory-nociceptive ‘danger detection’ system. Sixteen subjects executed a three-oddball auditory task, while receiving either painful or non-painful thermal stimulations. Speech vowels (standard, target) or white noise (distracter) of 60ms were presented through shielded insert earphones (Etymotic) at 1.4Hz (Slow task) or 8Hz (Fast task). Thermal stimuli were produced by a 3x3cm contact thermode (Medoc) applied to the forearm. Stimulation consisted in plateaus of 15s. After each thermal block, subjects rated intensity and unpleasantness of the sensation using a visual analog scale. Skin conductance response (SCR) was measured throughout the experiment. Using a 64-channel EEG system (Biosemi), with a sampling rate of 16kHz, auditory evoked potentials (AEPs) of several timescales were recorded: from auditory brainstem response (ABR) to the auditory P3a/b complex.

Results
The auditory task had a significant effect on pain ratings, with pain judged as less intense and unpleasant during the Slow task. A similar pattern was found in the SCR. Comparing the effect of painful and non-painful stimulations on AEPs revealed a distinct pattern of modulation, both in terms of temporal dynamics and topographic maps. In the presence of pain, early auditory responses are enhanced whereas later components such as the P3b are reduced.

Conclusion
By combining several methods and measuring AEPs at different timescales, we were able to assess both high-level and direct low-level interactions, between audition and nociception. Our results support the existence of a pain specific modulatory mechanism that affects early auditory processing. Results on auditory-nociceptive interactions also contribute to better characterize the human efferent system.

Effects of Unilateral Conductive Hearing Loss On Auditory Brainstem Responses in Rats
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Background
Listeners with unilateral conductive hearing loss (UCHL) have reduced their ability to localize sound. The effects of UCHL along the central auditory pathway are still poorly understood. Currently, most available information focuses on the ipsilateral effects of UCHL. However, due to the extensive crosstalk along the entire central auditory pathway, we hypothesize that UCHL will also affect the contralateral side of the central auditory pathway. Another important but still unresolved aspect is whether the effects of UCHL are reversible, indicating the existence of experience dependent plasticity in the central auditory pathway.

Methods
To test this we used an animal model of UCHL by earplugging. Earplugs were inserted in the right external canal of young adult Sprague-Dawley rats. Neuronal electrical activity was measured in both plugged (ipsilateral) and unplugged sides (contralateral) by auditory brainstem response (ABR) tests, in response to clicks and pure tones before, during (10 days), and 10 days after earplug removal.

Results
Our results from clicks in the plugged-side showed a threshold increase of 40 dB and smaller wave I, II and III amplitudes. In the unplugged-side, we found no change in the thresholds, but interestingly wave II and III amplitudes were significantly
smaller. The analyses of the latencies (peak and interpeak) showed an increase only in the plugged-side. We also analyzed the ratio of amplitude waves III/I. Data showed an increase in the ratio in the plugged-side, whereas there was a decrease in the unplugged-side. After earplug removal the threshold and latencies recovered in the ipsilateral side, whereas the amplitude was only recovered for waves II and III, not for wave I. In the contralateral-side the amplitude waves were fully recovered.

Conclusion
Our results suggest that UCHL alters reversibly and bilaterally the firing rates and/or synchrony of firing neurons in the auditory brainstem. The decreases in amplitude after UCHL might also reflect an imbalance between excitation and inhibition in the lower auditory brainstem.

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Impact of Kv3 Channel Modulator AUT3 on In Vivo Auditory Processing in Mice
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Background
Voltage-gated K+ channels of the Kv3 subfamily activate at strongly depolarised membrane potentials to drive quick repolarisation after an action potential, and also rapidly de-activate to enable fast re-initiation of action potentials. Kv3 channels are strongly expressed in the central auditory system, at levels that can be modulated by auditory experience (such as noise exposure). These channels are therefore potential targets for drug treatments aimed at ameliorating central auditory pathologies arising from noise exposure. The aim of this study was to determine whether a novel modulator of Kv3 channel activity, AUT3, alters auditory brainstem responses (ABRs) in normal mice or in mice exposed to mild acoustic trauma.

Methods
ABRs were recorded from anaesthetised naïve or noise-exposed CBA/Ca mice. Noise-exposed mice had been subjected to 105 dB SPL, 8-16kHz noise under anaesthesia, 1 day prior to ABR measurements. ABRs were recorded 30 minutes before and 2, 32, 62, 92 and 122 minutes after intraperitoneal administration of either AUT3 (60mg/ml) or vehicle solution. Auditory stimuli were presented free-field, and included 0-80 dB SPL clicks, as well as broadband noises followed by clicks. Analysis involved calculation of ABR thresholds, peak-to-trough wave amplitudes, wave peak latencies, inter-peak latencies and root-mean-square overall amplitudes.

Results
In both naïve and noise-exposed mice, there was a small but significant increase (p<0.02) in the inter-peak latency of ABR wave I to IV following AUT3 but not vehicle injections. The latency shift in wave IV was evident in absolute as well as inter-peak latency analysis, whilst the absolute latency of ABR wave I was not significantly altered. No significant effect of AUT3 injections (relative to vehicle injections) was observed for ABR thresholds, wave amplitudes, or overall root-mean-square amplitudes in either group of animals.

Conclusion
AUT3, a Kv3 channel modulator, caused a small but significant increase in the latency of ABR wave IV in both noise-exposed and naïve mice, but no detectable change in either the latency of ABR wave I, ABR thresholds, peak-to-trough wave amplitudes, or overall root-mean-square amplitudes. These findings indicate that AUT3 causes no gross abnormalities in the ABR, but does have a subtle effect on the timing of late ABR waves, which may reflect an impact on auditory processing in higher central auditory structures such as the inferior colliculus.
Musical training strengthens the subcortical-cortical encoding and categorical perception of speech
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Background
There is increasing evidence suggesting that musical training alters the psychophysiological processing of complex sound, including speech. Given that speech sounds are typically perceived in a categorical manner, we reasoned that musicianship might also enhance the categorization/classification of speech and corresponding phonemic-level neural representations.

Methods
To this end, we compared cortical and brainstem event-related potentials (ERPs) elicited by a speech sound continuum in musician and nonmusician listeners.

Results
Behaviorally, musicians obtained steeper identification functions and classified speech sounds more rapidly (i.e., shorter reaction times) than their nonmusician counterparts. Analysis of the underlying neuroelectric activity revealed that relative to nonmusicians, musicians showed more robust and coherent phase-locking to speech formant energy in brainstem ERPs coupled with complementary increased magnitude (N1-P2 component) in cortical evoked responses. Identification functions derived from cortical potentials showed that neuronal activity could accurately predict listeners’ perceptual boundary and phonetic classification of the speech continuum for both groups. While neural indices (e.g., magnitude) extracted from brainstem and cortical ERPs were correlated with behavioral measures across the board, these brain-behavior associations were generally stronger in musically trained individuals.

Conclusion
Results suggest that in addition to enhancing the overall salience of speech encoding at multiple tiers of the auditory pathway, musical training strengthens the coordination of processing between sensory and perceptual levels of the auditory brain. We infer that extensive musical training acts to refine a hierarchy of internalized representations for auditory objects thus supplying more faithful phonemic templates to the decision processes subserving speech sound identification.
Individual Differences in the Perception of Musical Consonance

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Background
A consonant sound is perceived as 'resolved', 'stable' and 'pleasant'. When two tones are played simultaneously (to produce a dyad), the frequency ratio between them determines the musical interval. Different intervals evoke differing degrees of consonance. Individual listeners have different preferences for consonant dyads over dissonant dyads. Individual preference for consonance correlates with preference for sounds with frequency components that closely fit a single harmonic series (McDermott et al 2010) suggesting that the perception of consonance may be related to the perception of harmonicity.

Neurons in the auditory pathway tend to synchronise their firing to periodic sounds, suggesting that pitch may be coded in the inter-spike intervals (ISIs) of neuronal firing patterns. Consonant musical intervals produce stronger representations of harmonically relevant periodicities in the ISI distribution of the auditory nerve than dissonant intervals, suggesting that this may be a sub-cortical mechanism contributing to the perception of consonance (Ebeling 2008; Bidelman and Krishnan 2009). It is therefore possible that individual differences in consonance correlate with individual differences in the neural representation of dyads.

Methods
We compared consonance ratings for both diotic and dichotic intervals to measures of the frequency-following response (FFR). The FFR is an electrophysiological measure of phase locking in the brainstem.

We used the procedure of McDermott et al (2010): individual preference for consonance for 19 normal hearing listeners with a range of musical experience was calculated by subtracting their average pleasantness rating of three dissonant dyads from their average rating of three consonant dyads. We then measured the FFR to a Perfect 5th musical dyad (a consonant interval) in order to assess individual sub-cortical temporal coding acuity. An autocorrelation function of each FFR was used to extract the strength of coding of stimulus periodicity.

Results
Individuals with a stronger preference for consonant musical dyads show a stronger representation of stimulus periodicity in the FFR for both diotic and dichotic dyads.

Conclusion
The results of this study demonstrate that more robust neural coding of pitch relevant periodicity at a subcortical level corresponds to a greater preference for consonance. The correlation was significant for both diotic and dichotic stimuli, providing support for models in which consonance is the consequence of increased neural synchronisation (Ebeling, 2008; Shapira Lots and Stone, 2008). The findings suggest that individual behavioural differences are driven in part by individual differences in how precisely populations of neurons are able to phase-lock to pitch-relevant periodicities.
Attention-Modulated Responses in the Cortex and Brainstem to Ongoing Speech Streams

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Background
Selective attention modulates neural activity recorded in the auditory cortex. However, little evidence exists that attention modulates activity in the auditory brainstem, despite the presence of efferent connections from auditory cortex to subcortical structures. To probe the effects of attention in both the cortex and the brainstem simultaneously, we recorded high-frequency EEG from listeners during a dichotic listening task using speech stimuli.

Methods
We recorded 32-channel EEG at 16,384 Hz from listeners selectively attending to one of two speech streams. Each speech stream consisted of a random string of digits (1-5). Listeners monitored one of the two streams for any occurrence of two back-to-back, consecutive digits within the ~60 s long stimuli. Onsets of digits within each stream were separated by 500 ms, and the mixture was presented such that the to-be-attended stream started 250 ms prior to the to-be-ignored stream (i.e., digits in the two streams alternated temporally, although they overlapped). To allow separation of the brainstem frequency following response (FFR) to each stream, stimuli were presented dichotically (one stream in each ear) and were each vocoded using pulse train carriers of differing repetition rates (97 and 113 Hz) and different frequency bands (so that each stream excited a distinct set of peripheral auditory filters). Half of all stimuli were reversed in polarity to enable analysis of the neural envelope and carrier FFRs. Responses to the onset of each digit were examined in the time domain for the cortical response, and in the frequency domain for the FFRs.

Results
Cortical responses to attended and ignored digits showed large differences at most scalp electrode sites, although response morphologies varied across subjects. Applying a combination of PCA and SVM resulted in classification of attended and ignored responses at above-chance levels when small numbers of individual responses were averaged together. In contrast, brainstem envelope FFRs exhibited phase locking to the fundamental frequency of each stream for all subjects, but there were no systemic differences in the FFR strength in response to the streams depending on how attention was directed. No phase locking was observed in the carrier FFRs.

Conclusion
Despite inter-subject differences in response morphology, cortical responses were clearly modulated by selective attention. No such effects were seen in the brainstem response despite the care taken in designing stimuli to maximize the separability of the subcortical responses to the two streams.

The Role of Subcortical Encoding in Accounting for Speech Perception in Steady-state and Amplitude-modulated Noise

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Background
One important factor that makes young normal-hearing listeners able to understand speech in a considerable amount of noise is the fact that they can ‘listen in the dips’ of time-varying maskers. This benefit has been hypothesised to depend...
upon accurate neural phase locking to the temporal fine structure to exploit periodicity in the target and/or masker, and upon the ability to track slower envelope fluctuations.

However, speech perception in noise not only depends on bottom-up auditory processing. The success of speech perception in both steady-state and amplitude-modulated noise also depends on higher order cognitive processes.

This study examines the relative contribution of both low level auditory processes and higher level cognitive processes to the perception of speech in noise. In particular, the role of subcortical encoding of speech in different types of maskers is assessed.

Methods
Auditory brainstem responses were measured in normal-hearing listeners to a steady-state vowel in quiet, steady-state speech-shaped noise (SSN), and amplitude-modulated speech-shaped noise (AMN). In addition, temporal processing abilities were assessed by measuring thresholds for gap detection, amplitude modulation detection, and frequency modulation detection. To assess the contribution of cognitive factors, measures of processing speed, attention, and working memory were also obtained. Furthermore, a visual analogue of the speech perception in noise task was conducted (Text Reception Threshold). Of primary interest is the extent to which the various measures, both low and high level, correlate with listeners’ abilities to perceive speech in SSN, AMN, and two-talker babble.

Results
Our pilot data indicate that voice pitch is less robustly encoded at the brainstem in background noise. Furthermore, in AMN the pitch may be more robustly encoded at the amplitude dip of the masker.

Conclusion
We expect to find that both low level auditory and higher level cognitive processes contribute to listeners’ abilities to understand speech in noise. However, we expect that the robustness of subcortical encoding of speech will be the primary predictor for people’s abilities to understand speech, especially in AMN. Subcortical encoding of speech is expected to be more robust in the dips of AMN. Furthermore, temporal processing abilities are expected to correlate highly with both subcortical encoding and speech understanding, particularly in AMN.
Hyperexcitability of Inferior Colliculus Neurons Caused by Acute Noise Exposure
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Background
Noise exposure is one of the most common causes of hearing loss. Recent studies found that noise exposure induced cochlear damage may change the excitability and tonotopic organization of the central auditory system (CAS). This plasticity was suspected to be related with tinnitus and hyperacusis. However, how cochlear damage affects CAS function and causes these neurologic diseases is still not clear. Since CAS function is activity dependent, we hypothesize that a restricted cochlear lesion might disrupt the balance of excitation and inhibition in the CAS, and therefore affect its neural activity.

Methods
Extracellular recording from the inferior colliculus (IC) neurons was recorded using a 16-channel microelectrode (NeuroNexus, A1x16-5mm-100-177) in anesthetized rats before and after the noise exposure. The multiunit discharges were recorded using OpenEx (TDT) software and the peri-stimulus time histograms (PSTH), frequency response areas (FRAs) and spiking rate level function were reconstructed using custom software. A narrow band noise (105 dB SPL, 20 kHz, 1000 Hz bandwidth) was presented by a high frequency speaker (FT28D) for 30 minutes to produce a restricted lesion in the cochlea.

Results
The noise exposure caused a dramatic decrease of the characteristic frequency (CF) in about 2/3 of the high frequency neurons with/without causing a significant threshold shift. The noise exposure also caused an increase in firing rate of the low frequency neurons at super-threshold levels whereas it dramatically decreased the firing rate of the high frequency neurons. These results demonstrated that the hyperexcitability caused by noise exposure only occurred in the IC neurons tuned to frequencies lower than the noise exposure frequency; whereas the CF shift and tuning curve widening mainly happened in IC neurons tuned above the noise exposure frequency.

Conclusion
Our results suggest that noise exposure caused different plastic changes in IC neurons in different frequency regions. The reduced peripheral input following noise exposure may affect both the excitatory and the inhibitory circuits, especially the side-band inhibition at the lower edge of the noise exposure frequency, and affects the excitability and CF of IC neurons.

Pharmacological activators and inhibitors of Kv3.1 potassium currents
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Background
Kv3 potassium channels are expressed in the auditory brainstem and in rapidly spiking neurons throughout the brain, where firing at high rates with high temporal accuracy is required for sensory processing. Kv3.1 channels typically activate at positive potentials and have a very rapid rate of activation and deactivation in response to transient depolarization.

Methods
We now report the first compounds to specifically modulate Kv3 channels. These compounds are imidazolidinedione derivatives, Compound 1 (AUT1) and Compound 2 (AUT2). We have performed whole cell patch clamp recordings of Kv3.1 channels.

Results
Using CHO cells stably expressing Kv3.1 channels, we have found that 10μM AUT1 shifts the voltage of activation of Kv3.1 currents towards negative potentials such that currents are activated around -20 mV and produce a 131% increase in current at test potentials of -10mV. Numerical simulations of the firing properties of auditory brainstem neurons, predict that increasing concentrations of AUT1 would be expected to decrease firing rate in response to high frequency stimulation (400 Hz), but to increase the temporal accuracy with which actions potentials are phase-locked to the stimuli. In contrast, 10μM AUT2 produced a sustained shift in voltage-dependence of inactivation to more negative potentials after three minutes of incubation, as well as altering voltage-dependence of activation. Although Kv3.1 channels usually inactivate only very slowly during sustained depolarization, the rate of channel inactivation is also markedly increased in
the presence of AUT2. Thus the net effect of this compound is to suppress Kv3.1 currents in the physiological range of membrane potentials. In numerical simulations, AUT2 had a biphasic effect on excitability. Low concentrations increased the rate of firing in response to 400 Hz stimulation whereas higher concentrations prevented neurons from responding to high-frequency stimulation, as is found in mice in which Kv3.1 has been deleted.

**Conclusion**

Pharmaceutical modulation of Kv3.1 currents represents a novel avenue for manipulation of neuronal excitability, and has the potential for therapeutic benefit in the treatment of hearing disorders associated with central auditory deficits.

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**Distribution of Vesicular Glutamate Transporters in the Cochlear Nucleus of C57 Mice**

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**Background**

The cochlear nucleus (CN) is a multisensory integration center in the auditory pathway, receiving innervation from auditory and somatosensory structures. Vesicular glutamate transporters, VGLUT1 and VGLUT2, have distinct distributions in the CN in guinea pigs and rats: VGLUT1 is highly expressed in the magnocellular area of the ventral CN (VCN), whereas VGLUT2 is predominantly expressed in the granular cell domain (GCD), which includes DCN2 and shell region of the VCN. The GCD receives non-auditory inputs from somatosensory nuclei, including spinal trigeminal nucleus (Sp5) and cuneate nucleus (Cu). VGLUT1 and VGLUT2 have been used as the excellent markers to examine auditory nerve degeneration and somatosensory compensation after unilateral deafness in guinea pigs (Zeng et al., J. Neuroscience, 2009). C57 mice are prime candidates for genetic mutations in hearing system. However, the distributions of VGLUT1 and VGLUT2 have not yet been reported.

**Methods**

Here we examined the distributions of VGLUT1 and VGLUT2 in C57 mice in comparison with that in guinea pigs. Cresyl violet nissl stain was used to confirm the different layers and cell types in the CN. Metamorph software was used for quantification of VGLUT1 and VGLUT2 immunoreactivity in each CN region.

**Results**

The cytoarchitecture in the CN of C57 mice is similar to that of guinea pigs except for a large interstitial region (INT) in anteroventral CN (AVCN) in the C57 mice. The most intense VGLUT1 labeling in C57 mice was in the molecular layer of dorsal CN (DCN), shell region and magnocellular area of ventral CN (VCN), in contrast to less intense VGLUT1 labeling in the shell regions of guinea pigs. Like the guinea pig, the most intense VGLUT2 labeling in C57 mice was in the GCD, but VGLUT2 in C57 mice was more intense than guinea pigs in DCN3 and VCN. Similar to guinea pig, VGLUT2 was more intense than VGLUT1 in the DCN2 and DCN3, whereas VGLUT1 was more intense than VGLUT2 in the VCN and DCN1.

**Conclusion**

When compared with guinea pigs, VGLUT1 and VGLUT2 in C57 mice shows similar distribution in the CN, however, some regions show different intensities. This study will lay the foundation for studies of VGLUT distribution changes after deafness.

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**Assessing the Relation Between Auditory Nerve Fibre Deafferentation, Hearing Threshold Increase, and Spontaneous Neuronal Hyperactivity**

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**Background**

A recent study in mice has shown that normal hearing thresholds do not necessarily indicate the absence of cochlear damage; following temporary threshold shifts induced by noise exposure, a permanent reduction in wave I of the auditory brainstem response (ABR) is observed, together with deafferentation of auditory nerve fibres (ANFs) [Kujawa and Liberman (2009), J. Neurosci 29:14077-14085]. A similar reduction of ABR wave I has been observed in tinnitus subjects with normal audiograms, also suggestive of ANF deafferentation [Schaette and McAlpine (2011), J. Neurosci 31:13452-
The amplitude of the centrally-generated wave V, however, was not reduced in the tinnitus group, suggesting a compensatory increase in neuronal response gain that might underlie the generation of a neuronal correlate of tinnitus.

**Methods**
CBA/Ca mice were exposed to an octave-band noise (8-16kHz) at either 96dB or 105dB SPL for 2 hours under anesthesia. ABRs were recorded pre-trauma, 1 day post-trauma, and 4 weeks post-trauma. Extracellular multiunit recordings from the inferior colliculus were obtained 4 weeks post-trauma. All recordings were performed under ketamine-medetomidine anaesthesia.

**Results**
For all trauma groups, ABR thresholds were significantly increased, and ABR wave amplitudes strongly reduced, 1 day post trauma. 4 weeks later, thresholds had recovered fully in the 96dB group, whereas the 105dB group still showed threshold elevations. The amplitudes of ABR wave I remained decreased in both groups, suggesting deafferentation of ANFs. Amplitudes of ABR waves II-IV remained below pre-trauma values in the 96dB group. However, in the 105 dB group, amplitudes of wave IV recovered and reached pre-trauma levels at high sound intensities, suggesting a compensatory increase in neuronal response gain.

Extracellular multiunit recordings were obtained from the inferior colliculus 4 weeks post trauma. In the 96dB group, there was no significant increase in spontaneous firing rates. However, in the 105dB group, spontaneous firing rates showed a significant increase for neurons with tuning in the trauma frequency range, which was correlated with increased response thresholds.

**Conclusion**
The development of spontaneous neuronal hyperactivity in the inferior colliculus seems to require a minimum level of cochlear damage, demonstrating a non-linear relation between damage, plasticity, and spontaneous neuronal activity.

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Temporal coherence patterns in auditory cortex neuronal activity during an auditory streaming paradigm
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Background
Auditory scene analysis refers to grouping or segregating features of a complex acoustic input in order to identify and process the underlying objects. Auditory streaming is a simplified version of this problem, where two alternating rhythmic stimuli are perceived as two separate “streams” or fused into one, depending on a number of defined factors. However, the physiological mechanisms underlying streaming are not understood. We tested the hypothesis that objects (i.e. streams) may be represented in auditory cortex by neural assemblies with temporally coherent activity patterns, whereas multiple objects may arise from multiple non-coherent assemblies.

Methods
We employed voltage-sensitive dye imaging to record ongoing subthreshold neural activity across multiple fields of the auditory cortex in anesthetized guinea pigs. Imaging was performed at a 250 Hz sampling rate with 100x100 imaging pixels that covered 10x10 mm of cortex. Each experiment began with measuring responses to pure-tone stimuli ranging from 0.5 - 32 kHz, to define the best-frequency of individual pixels and the full tonotopic organization of cortex.

Results
We found that in the presence of a single rhythmic (4 Hz) pure-tone stimulus, membrane potentials from regions representing that stimulus reliably entrained to the rhythm. The phase coherence of this 4 Hz activity was increased relative to a silence condition, across a 4-octave region of cortex. We next addressed temporal coherence patterns in the presence of two out-of-phase rhythmic stimulus trains of pure-tones, as a function of their frequency difference (deltaF, from 1 to 18 semi-tones). We found that temporal coherence was largely dependent on deltaF: as deltaF increased, the two tones were represented by increasingly distinct (non-coherent) neural assemblies within individual areas. Temporal coherence did not depend on spatial segregation in cortex, as high coherence was also observed between distant subregions from separate cortical areas that shared best-frequency tuning. Finally, even at relatively large deltaF when temporally-distinct neural assemblies were present, some cortical regions were still responsive to both tones.

Conclusion
Our findings support a model whereby bottom-up object segregation may be enhanced by the formation of temporally incoherent neural assemblies. More specifically, the probability that two features will be grouped or segregated may vary with the degree of temporal coherence between neural assemblies representing those features, and intermediate values of coherence may be permissive for bistability.
Switching Auditory Attention Using Spatial and Non-spatial Features Recruits Different Cortical Networks

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Background
Attention is a selective process that allows us to tune into one of potentially several possible sound sources of interest. In vision, the ability to flexibly redirect attention based on task goals and stimulus properties has been shown to involve a distributed cortical network including the right temporal-parietal junction (RTPJ), as well as multiple frontal areas including the frontal eye fields, which are involved both in gaze control and deploying attention. However, the cortical network controlling auditory attention has been studied less, and in particular, how attention might be directed to non-spatial stimulus features (such as pitch) is not clear.

Methods
Subjects performed a task where they counted the occurrences of a specific letter (“E”) from one of two simultaneous talkers, each saying streams of six consecutive letters. Talkers were either spatially separated with the same pitch, or spatially collocated with different pitches. Subjects were cued at the beginning of the trial to attend to one talker (left, right, high pitch, or low pitch) for either the whole trial (maintain-attention condition) or for only the first half of the trial, switching to the other talker after three letters for the remainder of the trial (during a 600 ms gap period; switch-attention condition). Using magneto- and electro-encephalography combined with structural MRI information, we performed a within-subjects contrast of cortical activation when subjects maintained versus switched attention on both space and pitch trials.

Results
When switching versus maintaining auditory attention, using spatial stimulus features more strongly engaged RTPJ, consistent with previous results. However, when using non-spatial stimulus features, a distributed network including left auditory cortex and right frontal areas was more active.

Conclusion
Switching attention using spatial and non-spatial features differentially engages separate cortical networks. This suggests that top-down control of auditory attention involves the deployment of different processing strategies based on the stimulus features available for stream segregation.

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Induced gamma-band synchronization for auditory stream segregation in auditory cortex of the awake rat

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Background
Perceptual segregation of alternating tone sequence differing in frequency (ABA-ABA-...) depends on the frequency differences ($\triangle F_s$) between A and B tones and the inter-tone intervals (ITIs) between successive tones as indicated in the van Noorden's perceptual boundary. In the auditory cortex, tonotopic separation and forward suppression has been considered as possible neural underpinnings of this psychophysical phenomenon, which, however, cannot completely explain the phenomenon. Recent studies have proposed that synchronized ensemble activities contribute to formation of perceptual objects. To investigate whether such processing is also critical for auditory streaming, multichannel recording methods are effective which meet requirements of both covering a targeted functional structure and achieving higher spatial resolution capable of evaluating correlation between neural ensembles in each functional structure. Secondly, it is important to examine how synchronized or desynchronized activities based on functional structure contribute to stream formation.

Methods
The present study developed a method that can acquire multiunit activities (MUAs) and local field potentials (LFPs) from auditory cortex of the awake rats with microelectrode arrays capable of covering auditory cortical fields with relatively high spatial sampling resolution. We investigated the relationship between oscillatory synchronization of LFPs or spike - LFP phase coupling and acoustic condition in ABA-tone sequence.

Results
Response magnitudes such as spike rate of MUA and amplitudes of raw / gamma-band LFP to A tone increased with larger $\triangle F$ and longer ITI at recording sites selective to A-tone frequency, suggesting that tonotopic separation and forward suppression was most effective at the sites and not a correlate with the perceptual boundary. On the other hand, desynchronization across auditory cortical fields but tone selective synchrony was enhanced compared to those under anesthetized condition. Furthermore, different from evoked gamma-band oscillation, synchrony and power of induced gamma-band oscillation tended to be enhanced across neuronal ensembles selective to A tone under shorter ITI and larger $\triangle F$. Especially spike-phase coupling significantly increased under those conditions, suggesting that these neuronal characteristics were closely correlated with the perceptual boundary.

Conclusion
Not transient but phase synchrony, spike-phase coupling or sustained power of induced gamma-band oscillation are possible neuronal representations of the perceptual boundary of auditory streaming.
Re-Examining the Evidence for a Neural Representation of Pitch in Human Auditory Cortex
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Background
A number of functional criteria have been proposed for defining a human pitch center including: (1) greater activation for a pitch-evoking stimulus than for a matched control, and (2) greater activation for a strong pitch percept than a weak one. Some human investigations using functional magnetic resonance imaging attribute such patterns of pitch-related activity to lateral Heschl’s gyrus, but not all. In particular, Pena-gos and colleagues (2004) report sensitivity to pitch salience in this region, but used a small sample size (n = 5) that precluded group statistics.

Methods
The current replication reports a larger sample size, allowing group-level analysis and generalizability of results. Eighteen listeners (right-handed; 9 females) were scanned during passive listening. Four harmonic-complex tone conditions (3 strong in pitch salience, 1 weak in pitch salience) were presented, plus two spectrally matched ‘no pitch’ noise controls and a silent baseline condition. Signals were presented in a noise background to reduce cochlear distortions. Each sound condition was a 32-s dynamic sequence where the pitch varied within a given narrow range. Image data were analysed using SPM8 and subjected to random-effects group analysis, incidence mapping and region-of-interest analysis.

Results
In the group analysis, pitch- and salience-related activity were defined by the contrasts ‘pitch>noise’ and ‘strong>weak’, respectively. Both these contrasts showed differential activity which was reliable across the group (p<0.001, uncorrected) and widespread across auditory cortex. However incidence mapping revealed that this spatial distribution was not consistent in every listener (individual level p<0.01, uncorrected), which highlights the risks of generalising from a few participants. Region-of-interest analysis confirmed that the resolved complex tone conditions (strong pitch) elicited much greater activity than unresolved (weak pitch) and matched noise controls. Although the response to the unresolved complex>noise was small, the incidence map did show that a differential effect was present in some listeners, thus confirming some degree of sensitivity to pitch salience within these regions. Interestingly, planum polare showed a significant right hemisphere preference for resolved complexes, which was not observed for the other sound conditions or brain regions.

Conclusion
Our findings broadly support Penagos, but extend current knowledge. (1) Neural representations of both pitch and pitch salience are co-localized within multiple regions of human auditory cortex. (2) Presenting pitch signals as a random melody sequence promotes right hemispheric involvement in planum polare. Not only do these results support pitch coding in auditory cortex, but also provide further evidence for a processing hierarchy.
Predicting the Attended Speech Stream from the Ongoing EEG Response in a Cocktail Party Paradigm
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Background
In this study we aim to identify which of two ongoing speech streams a listener is attending using electroencephalography (EEG). The majority of work done in classification of auditory attention using EEG has used regular sequences of simple stimuli like noise or tone bursts, which evoke clear onset responses that are modulated by attention. We classify the attended stimulus by correlating the EEG response to the known envelopes of the speech stimuli using time-lagged canonical correlation analysis (CCA).

Methods
Data were collected for a previous study [Power et al., Eur. J. Neurosci. 35(9), 2012] in which EEG was recorded while subjects attended to one of two audiobooks, presented dichotically (one to each ear) at equal intensities. EEG was recorded using a 128-channel cap (downsampled to 32) with a BioSemi ActiveTwo system, at a sampling rate of 512Hz. The stimuli were presented in approximately thirty 60s blocks. Subjects had to answer questions about both stories to confirm that they attended to the desired one. We projected out blink artifacts in the EEG using the signal space projection method and applied a high pass filter at 1Hz. We then performed CCA at all possible time lags on the training data, and picked the lag and spatial weighting that yielded the maximum canonical correlation with the speech envelope. The test data were then weighted with the spatial weights and correlated with the envelope of the two, competing speech streams in the test block at the optimal lag. The stimulus with the largest correlation value was classified as the attended speech stream. Performance for each of three classifiers was compared: trained to the attended stream, the unattended stream, or the sum of the two streams. A leave-one-out cross validation was used on the blocks.

Results
Preliminary results indicate that training the canonical correlation classifiers at longer lags (between 160-240ms) yields a more accurate classification of the attended stream (82.21% +/- 4.71% standard error across subjects) than at shorter lags in the 60-140ms range (65.57% +/- 4.99% standard error across subjects).

Conclusion
Prior EEG studies that classify which of multiple concurrent stream is being attended by a subject use attentional modulation of the N100 (which occurs 100ms after stimulus presentation). We show that for continuous speech, later evoked potentials (around 200ms) are better at differentiating which stream a listener attended.
Exploring the perception and neural representation of vowels in ferrets
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Background
Object identification depends upon the exploitation of variation in object properties. In speech, vowels can be distinguished on the basis of formant frequencies – peaks in the spectral envelope introduced during articulation. Specifically, the location of first (F1) and second formants (F2) and distance between formants (F2-F1) offer cues for vowel identification across different talkers and voice pitches [Peterson GE, Barney HL (1952), Kewley-Port D, et al. (1996)]. Vowel identification can be studied at behavioral and neuronal levels using animal models such as the ferret: Ferrets can identify vowels differing in F1 and F2 and the spectral envelope of such vowels modulate the responses of neurons across ferret auditory cortex [Bizley JK, et al. (2009); Walker KM, et al. (2011); Bizley JK et al. (In Press)].

Methods
Here, we describe results from behavioral and neurophysiological investigations of vowel discrimination in ferrets. In our behavioral study, ferrets (n = 4) were trained to associate water rewards at different locations with two artificial vowels that differed in F1 and F2 positions. Subjects were then presented with probe trials in which the F1, F2 or F2-F1 of training vowels were swapped. In our neurophysiological study, the same stimulus set of vowels and probes were presented to untrained anesthetized ferrets (n = 3) whilst multi-unit responses were recorded from auditory cortex.

Results
Trained ferrets responded to mismatch vowels more frequently at the location previously associated with F2 than F1. Discriminability (d') between vowel pairs was also positively correlated with F2 separation between vowels. Neither F1 separation nor F2-F1 separation was correlated with discriminability. Neural discriminability was measured as spike distance between PSTH responses to vowel pairs. Spike distance was well correlated with F1, F2 or F2-F1 separation between vowels in a subpopulation of vowel-sensitive units (29/84; 35%). However in the majority of vowel-sensitive units (55/84; 65%), discrimination was only weakly correlated with cue separation.

Conclusion
These findings suggest that F2 position may disproportionately influence the perception and discriminability of vowels in ferrets. The underlying physiological basis for such behavior may involve the contributions of cue separation sensitive units observed here; however this will require further examination in trained, behaving animals.
Electrophysiology and perception of speech in noise: effects of age and hearing impairment
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Background
A common complaint of individuals with hearing impairment and older individuals is difficulty understanding speech in background noise. Despite being such a common problem, perception in noise difficulties are often not addressed clinically, leaving the affected population to suffer with decreased communication, additional stress, decreased participation in social activities, and poorer quality of life. Our approach is to use a combination of brain and behavioral measures to better our understanding of performance variability across individuals as a means of improving diagnosis and individualizing treatment. By measuring the ability of the brain to encode speech in noise we can begin to determine what neural information is available to the listener to ultimately understand speech in a noisy environment. Recent electrophysiological studies demonstrate that signal-to-noise ratio (SNR) is an important factor contributing to the timing and amplitude of cortical auditory evoked potentials (CAEPs) in young, normal-hearing individuals. The purpose of this study was to extend those previous CAEP findings, by testing older, hearing-impaired individuals and to explore the relationship between CAEPs and behavioral responses.

Methods
CAEPs and behavior measures were collected from 15 younger normal-hearing individuals, 15 older normal-hearing individuals, and 15 older hearing-impaired individuals using signal-in-noise stimuli. CAEP measures included the passively elicited P1-N1-P2 complex, which was elicited by a 1000-Hz tone and the syllable /ba/. Behavioral measures included sentence-level and word-level stimuli presented in noise. Target signals were presented at four levels (50-80 dB in 10 dB increments) in steady-state speech spectrum noise that was also varied in level (40 to 90 dB) resulting in signal-to-noise ratios of -10 to 35 dB.

Results
Results from these studies demonstrate that key factors affecting neural coding and perception of speech in noise include signal type, signal-to-noise ratio, age, and hearing status. In addition, more robust and synchronous neural coding is associated with better understanding in noise. Significant correlations were found between CAEP measures and perception-in-noise measures. Differences in sensitivity between the two measures will also be discussed.

Conclusion
Further examination of the important factors affecting brain and behavior and investigation of the relationship between these two measures will improve our understanding of speech perception-in-noise difficulties and help us to improve diagnosis and treatment of individuals with these problems. [Research supported by VA/RR&D (C6971M, C4844C) and the NIH/NIDCD (R03DC010914)]
The effect of attention and temporal shift on the build-up of auditory streaming: A concurrent EEG and subjective study

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Background
Auditory streaming tends to “build-up” over time with a tendency for listeners to perceive two streams increasing with sequence duration (Anstis and Saida, 1985). In addition, imposing a frequency or temporal shift mid-ABA sequence may influence stream segregation (Chakalov et al., 2012). The aim of the present study was to obtain a measure of the disruption to auditory streaming build-up by imposing a temporal shift near the beginning, or close to the end of, an ABA-sequence under different conditions of attention. The magnetoencephalographic negative wave N1m may be used as an indicator of auditory stream segregation (Gutschalk et al., 2005). When the repeating sequence of a pure-tone triplet (ABA-) is perceived as a single stream, the N1m amplitude is relatively small and consistent with the inter-stimulus-interval (ISI) between successive tones, when the sequence is perceived as segregated, the N1m amplitude is relatively large and consistent with the longer ISI between tones of each segregated stream. The present study used the electroencephalographic (EEG) equivalent, wave N1 to investigate stream segregation using a combined EEG and psychophysical test approach.

Methods
The stimuli were 9-sec sequences of repeating ABA triplets, where A= 1260 Hz, B=1000 Hz and Δf = 4 semitones. A temporal shift of 40 msec was imposed on the B tone in either the 4th triplet (early condition) or the 11th triplet (late condition). The control condition had no temporal shift imposed on the B tone. For each temporal shift condition, participants were instructed to focus on an integrated or segregated percept. Participants responded by button-press to indicate if one or two streams were perceived whilst EEG recordings were obtained using a 32-electrode array. Fifteen participants participated and EEG and subjective responses were recorded concurrently.

Results
Results indicate a difference in the N1 amplitude between the attention conditions, when the temporal shift is imposed early, rather than later, in the ABA sequence.

Conclusion
The EEG and subjective responses appear to concur; allocating attention to either an integrated or segregated percept affects the detection of a temporal shift imposed on and ABA sequence, particularly so early on in the sequence, rather than later in the sequence.
A Computational Model of Spatial Tuning in the Auditory Cortex in Response to Competing Sound Sources

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Background
To better understand sound localization in the auditory system, much work has been done on the spatial tuning of single neurons along the auditory pathway. It has been established that single neurons in the midbrain are sharply tuned to preferred directions, while cortical neurons show much broader tuning. A recent experiment on cortical responses in birds found evidence of spatial tuning to multiple competing sounds in the cortex (Maddox RK et al. Competing Sound Sources Reveal Spatial Effects in Cortical Processing. PLoS Biol. 2012). This study described situations in which cortical neurons show broad tuning to single location stimuli, but develop spatial preference in response to a second competing noise source. In the current study, we construct a computational model to explore how a cortical network can utilize sharp spatial tuning in its inputs to extract the locations of competing stimuli in a noisy environment.

Methods
The model cortical network receives inputs from spatially tuned neurons from an input layer. The spike rates of the input layer neurons are implemented using cross-correlation and time delay methods, followed by a spike generator. A mid-layer of interneurons receives inputs from input layer neurons, and synapses onto cortical neurons. Both interneurons and cortical neurons are implemented using integrate-and-fire models. The time course and magnitude of excitatory postsynaptic currents (EPSC) and inhibitory postsynaptic currents (IPSC) at the synapses were varied. The output of the model is the response of the cortical neurons to competing sounds.

Results
The model demonstrates broad tuning to single sounds, and develops spatial preference when a second competing noise is presented. Spatial tuning to masked stimuli is achieved by the suppression of neurons tuned to non-preferred directions by neurons tuned to preferred directions. The slower decay in IPSC relative to EPSC is important in order for response to the preferred direction to take dominance.

Conclusion
This study proposes a cortical mechanism of sound localization in noise. The model suggests that, in noisy environments, multi-neuron networks may be able “tune in” to their preferred source. This may be the functional basis of spatial location and noise separation in the auditory system, and could be a key component in understanding more complex behavioral phenomena such as the cocktail party effect.

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Cortical Pitch Regions Respond Primarily to Resolved Harmonics Once Distortion Products are Masked

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Background
Pitch is a defining perceptual property of many real-world sounds. Classically, theories of pitch perception have differentiated between temporal and spectral cues. These cues are rendered distinct by the frequency resolution of the ear, such that some frequencies produce “resolved” peaks of excitation in the cochlea, while others are “unresolved”, providing a pitch cue only via their temporal fluctuations. Behaviorally, unresolved harmonics on their own produce a weak pitch relative to resolved harmonics. Human neuroimaging studies, in contrast, have often reported strong responses to the temporal cues conveyed by unresolved pitch stimuli. One possibility is that many of these studies were influenced by ‘distortion products’ - frequency components introduced by nonlinearities in the ear that can effectively act as resolved harmonics. Here we use fMRI to test the relative contribution of resolved harmonics, unresolved harmonics, and distortion products to the response of cortical regions sensitive to pitch.

Methods
We first searched for brain regions in each subject that responded more to pitch stimuli compared with noise, irrespective of resolvability, and we then compared responses to resolved and unresolved harmonics using independent data from
these same regions. Every condition was scanned with and without low-frequency noise designed to mask distortion products. This design allowed us to test whether masking noise enhances response differences between resolved and unresolved harmonics, as would be expected if distortion products generated by unresolved harmonics are effectively treated as resolved frequency components.

**Results**

We report: 1) Consistent with previous reports, all 23 subjects tested exhibited pitch-sensitive regions that responded more to harmonic tones than noise; these regions were predominantly localized to low-frequency and non-tonotopic regions of anterior auditory cortex. 2) These regions were driven primarily by spectrally resolved harmonics, although they also produced a weak but consistent response to unresolved harmonics relative to frequency-matched noise. 3) Notably, this response preference for resolved harmonics was only observed with concurrent noise designed to mask distortion products. Without masking noise, we observed strong responses to harmonic tones irrespective of resolvability, suggesting that distortion products have a surprisingly large effect on the cortical response, effectively acting as resolved frequency components and potentially explaining previously observed responses to pitch in the absence of spectral cues.

**Conclusion**

Our results suggest that resolved spectral information is critical to the response of pitch-sensitive brain regions and that resolved distortion products substantially enhance cortical responses to unresolved pitch stimuli when not masked.

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**Neural Correlate of Time Shift Detection in an ABA- Streaming Paradigm in the European Starling**

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**Background**

Segregating or integrating sounds from multiple sources is a crucial task of the auditory system for survival. For studying the mechanism of auditory scene analysis, sequences of two alternating tones are commonly used as stimuli (ABA-, A and B being different tones, - indicating a longer temporal gap). Using such stimuli, van Noorden (1975) reported that the temporal shift of the middle B tone in ABA- triplet was more easily detected when the consecutive sounds were perceived as one stream (fusion) compared to the situation where the sounds were heard as two streams (fission). The European starling (Sturnus vulgaris) showed a similar ability to detect the time shift of middle tones in a triplet depending on the frequency separation between consecutive tones suggesting a similar perception of stream segregation (Itatani, Cordes and Klump 2011). The change in neuronal representation in relation to fission or fusion, however, is still unclear. Here we combined the behavioral and physiological observations using the European starling to investigate the neural correlate of perceptual stream segregation.

**Methods**

Three European starlings were trained to respond to the time shift of B tones in ABA- tone triplets in an operant Go-NoGo task. Platinum-iridium electrodes were chronically implanted into the field L complex being homologous to the mammalian primary auditory cortex. During the behavioral experiment, spike responses were recorded using a radio telemetry system. The frequencies of the A and B tones were set so that their frequency separation was 0, 6 or 12 semitones. The amount of the time shift of the B tone was varied between 10 and 90%. For constructing neurometric functions, spike responses in each time shift condition were compared.

**Results**

The ongoing data collection suggests similar patterns of change of the d’ values of the psychometric and neurometric functions, which increased as the amount of the time shift increased. In general, the d’ values in neurometric function dropped if the frequency separation between two tones was increased.

**Conclusion**

The results suggest that in the European starling the representation of time shifts by the spike responses for different frequency separations of the tones are correlated with the behavioral observation of the accuracy of detecting a time shift.
Neuronal Representation of Temporal Regularity Associated with Pitch Perception in Macaque Auditory Cortex

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Background
Pitch is an auditory percept that requires neural representations of both stimulus regularity and perceived pitch. Recent human studies using direct recordings of local field potentials (LFPs) in the auditory cortex along Heschl's gyrus show that time-locked responses correlate with the temporal regularity of the stimulus, whereas sustained gamma oscillations correlate with perceived pitch (Griffiths et al., 2010; Sedley et al., 2012). The present study in a primate model system examined temporal regularity associated with pitch at both the single-unit level and at the level of cell assemblies (LFPs).

Methods
Stimuli contained a transition between noise and either a noise with a regular repetitive structure in time, known as a regular interval noise (RIN), or a harmonic complex, at seven different rates (i.e., 8, 16, 32, 64, 128, 256, and 512 Hz). Comparing neuronal responses above and below the lower limit of pitch (approximately 30 Hz) allows differentiation of responses related to pitch as opposed to stimulus regularity. LFPs were analysed for evoked potentials and time-frequency analysis.

Results
Single-unit activity (116 neurons) and LFPs (82 sites) were recorded simultaneously from the core and the lateral belt (LB) of macaque auditory cortex identified using fMRI. Evoked responses time-locked to the stimulus regularity were observed both below and above the lower limit of pitch, whereas induced, non-time locked, high-gamma (> 70 Hz) responses associated with the transition from noise to RIN were observed particularly for rates above the lower limit of pitch. Neuronal tuning specific to the different rates of the RIN and harmonic complex stimuli was compared at the level of single-units and LFP. The tuning of single units was sharper compared to LFP, yet both types of neuronal responses exhibited tuning above the lower limit of pitch, associated with pitch perception. There were some stimulus specific differences: (i) the magnitude of responses was stronger to the RIN than to the harmonic complexes, and (ii) responses to the RIN were sustained compared to the transient nature of response to the harmonic complexes.

Conclusion
The results seem to be topographically distributed in that the stimulus-regularity related responses were observed mainly in the AI, whereas the induced high-gamma responses related to the pitch percept were observed lateral to this in the belt. These results complement those from the recordings of LFPs in humans and provide insights into how single neuronal responses might support pitch perception.
MEG correlates of pitch perception in human auditory cortex
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Background
Despite its ubiquity in speech and music, the cortical origins and neural mechanisms underlying pitch perception are still not well understood. While some researchers have shown cortical correlates of pitch in lateral Heschl's gyrus (HG) (Patterson et al., 2002; Penagos et al., 2005; Pushmann et al., 2010), others argue for a locus in planumtemporale (PT) (Hall & Plack, 2009). A more distributed representation of pitch along the HG has also been argued (Griffiths et al., 2010), where different parts of HG have specific functions within a pitch system (Kumar et al., 2011). The predicted properties of such a 'pitch centre' (or higher-order 'predictive centre' in the pitch system) are that a) activity should be similar across different pitch types, e.g. harmonic complex (HC), clicktrain (CT), and regular-interval noise (RIN); and b) for each pitch type, activity will occur for a transition between a spectrally matched control stimulus (noise) and a pitch stimulus.

Methods
We used three different pitch types (HC, CT, RIN) and two different regularity rates (20 and 250 Hz), one below and one above the lower limit of pitch (Krumbholz et al., 2000). The stimuli used unresolved harmonics (including masking noise below the lowest harmonic) and spectra were 1-over-f shaped with a range between 15-3000 Hz. Participants listened passively to contiguous segments of either regular-noise-regular or noise-regular-noise stimuli.

Results
The results revealed a strong transition response from noise to pitch (250 Hz) for all three pitch types with a latency of ~130ms and an N100m-like topography. The opposite transition resulted in a P50m-N100m complex for transitions from HC and CT, while transitions from RIN did not yield any evoked response (see also Lütkenhöner et al., 2011). In contrast, transitions from noise to 20Hz regularity were generally much weaker (HC, CT) and non-existent for transitions to RIN. The opposite transition again resulted in a weaker P50m-N100m complex (HC, CT), with the exception of transitions from RIN.

Conclusion
The results confirm both of the predicted properties in that the response is similar for all three different pitch types used here and is specific to that transition, i.e. is different from the reverse transition. Analyses of the time-frequency response and beamforming algorithms are ongoing and will shed light on the cortical origins of these responses.

Cortical Pitch Response: Differential sensitivity to pitch contour and pitch direction
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Background
Voice pitch is an important information-bearing component of language and music that is subject to experience-dependent plasticity at both cortical and subcortical stages of processing. The components of the scalp-recorded cortical pitch response (CPR), reflecting synchronized neural activity relevant to pitch in the lateral Heschl's gyrus (the presumed site of pitch processing), may provide a new and complementary window to examine experience-dependent enhancements in pitch encoding previously observed in musicians and tone language speakers at the level of the brainstem and to shed more light on the hierarchical nature of pitch processing along the auditory pathways. We recently demonstrated that the CPR is sensitive to both pitch and its salience. Here we examine the sensitivity of the CPR components to changes in pitch contour, and pitch direction that are characteristic of lexical mandarin tones.

Methods
Cortical pitch (CPR) responses were evoked from 10 Chinese listeners in response to three IRN stimuli with: (1) a flat steady state pitch (SS); (2) a rising pitch contour approximating Mandarin lexical tone 2 (T2); and (3) a falling pitch contour approximating Mandarin lexical tone 4 (T4). Pitch specific response components were isolated by preceding the 250 ms IRN (evoking a pitch response) stimuli by a 500 ms noise precursor (evoking only the obligatory onset responses). Stimuli were presented binaurally at 80 dB SPL at a repetition rate of 0.93/sec.

Results
CPR response amplitude systematically varied with pitch contour and pitch direction with T2 eliciting the largest pitch response followed by T4, and the steady state stimulus eliciting the smallest CPR component with poorly defined response morphology. In addition, response to T4 exhibited the shortest latency followed by T2, with the response latency
for the steady state stimulus being the longest. These results suggest that the CPR is differentially sensitive to both pitch contour and pitch direction.

Conclusion
Results are consistent with the notion that the CPR reflects neural activity relevant to pitch that is differentially sensitive to certain pitch attributes (steady state vs. dynamic pitch; and rising vs. falling contour of pitch). Thus, the CPR, along with the brainstem frequency following response, provides a physiologic window to evaluate the hierarchical organization of early sensory level pitch processing at the cortical and brainstem levels that is subject to experience-dependent plasticity.

Functional NIRS Measurements of Auditory, Visual and Somatosensory Responses in Normal-Hearing Individuals
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Background
Deafness induces crossmodal reorganisation within the auditory cortex. Electrophysiology and functional neuroimaging studies suggest a link between cortical crossmodal activation patterns and cochlear implant performance. However, electrical artefacts and implant magnets confound EEG and fMRI recordings respectively. Near-infrared spectroscopy (NIRS) is suited to auditory neuroimaging following cochlear implantation since it is unaffected by electrical artefacts and the implant magnet. NIRS has been used successfully to assess auditory-evoked cortical responses in deaf children with cochlear implants. We present preliminary NIRS data from a study of multisensory responses (visual, auditory and somatosensory) in 10 normal-hearing adult participants (mean age 33±9 years, n=4 male).

Methods
Auditory stimuli consisted of sinusoidal amplitude-modulated (10Hz; 100% modulated) or unmodulated broadband noise at 80dB SPL. Visual stimuli consisted of 1000 white dots on a black background that were randomly flashing or coherently moving. Somatosensory stimuli were delivered to the palms of both hands via a custom-made vibrotactile stimulator. These consisted of 10Hz or 20Hz sinusoidal vibrations. Auditory, visual and somatosensory stimuli of duration 20s were presented five times each in a pseudorandom order, interleaved with rest periods (20s). NIRS data were acquired using a 24-channel Hitachi ETG4000 system. All data were preprocessed and statistically analysed using NIRS-SPM software and general-linear-modelling techniques.

Results
Cortical responses were successfully recorded in auditory (n=8), visual (n=9) and somatosensory (n=6) modalities. HbT changes in the temporal lobe, occipital lobe and post-central gyrus from a representative subject are shown (Fig1).
Auditory stimuli: 6 of 8 subjects exhibited areas in the temporal lobe with significantly greater HbT responses to modulated, compared with unmodulated noise. Visual stimuli: 6 of 9 participants showed areas in the occipital lobe with greater cortical response to coherent-moving, compared with random dots. Somatosensory stimuli: 20Hz vibration, compared with 10Hz, produced significantly greater responses in the region of the postcentral gyrus in 3 of 6 participants. Statistics significant to p<0.05; expected EC correction.

Conclusion
We successfully recorded cortical responses in auditory, visual and somatosensory modalities. Our future studies will use a similar paradigm to assess crossmodal activity in the auditory cortex of profoundly deaf adult subjects with and without a cochlear implant. We hypothesise that crossmodal activity measured in deaf individuals using NIRS may provide a useful prognostic indicator of performance following cochlear implantation.

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Crossmodal Influences on Steady-State Responses in a Subjective Audio-Visual Binding Task
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Background
When faced with a crowded auditory scene, such as the canonical cocktail party, speech can be better understood when a listener is able to see the talker's face. In this situation, the talker's voice and moving mouth vary coherently in time and are often perceived as "bound" together into one crossmodal object, an important phenomenon for understanding the sensory world. Much of the behavioral work examining crossmodal effects has focused on illusions, such as the McGurk effect. Electrophysiology has shown a number of brain regions to respond to both auditory and visual stimuli, and a subset of those regions show crossmodal interactions. Here we investigated binding based crossmodal temporal correlations, using human neuroimaging and stimuli created to share some of speech's properties, but to be free of the linguistic confounds of the cocktail party.

Methods
Neural activity was recorded using magneto- and electro-encephalography (M-EEG). Auditory stimuli consisted of an amplitude-modulated (AM) tone complex; visual stimuli consisted of a radius-modulated disc on a black background. The modulation envelopes were created by low-passing noise at 6 Hz. On each trial, the tone amplitude and the disc radius changed in one of three temporal relationships: larger amplitude, larger radius (0-phase); larger amplitude, smaller radius (pi-phase), and no relationship (ind-phase). Listeners were asked to answer the question, "Did the visual and auditory stimuli change as one?" in order to determine if binding was perceived. Additionally, auditory stimuli were "tagged" by applying sinusoidal AM at 37 Hz and visual stimuli were tagged by sinusoidally modulating the intensity of the center of the disc at 10 Hz, facilitating steady-state analysis (e.g., phase-locking value, PLV). Anatomically-constrained inverse imaging was used to localize data on the cortical surface.

Results
Listeners perceived binding much more frequently in the 0-phase and pi-phase conditions than in the ind-phase condition. The 0-phase condition was reported to bind slightly more often than the pi-phase condition. Additionally, some cortical regions showed significant steady-state differences between bound and unbound conditions.

Conclusion
It is possible to create stimuli for which listeners report strong binding. By tagging different modalities with orthogonal frequencies and using inverse imaging, we are able to create a cortical map of the crossmodal effects of binding.

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Cross-modal Alterations to Receptive Field Properties of the Auditory Cortex
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Background
Sensory integration occurs across various sensory systems and is observed in primary sensory cortices. Such cross-modal inputs might provide substrates for cross-modal compensation in the event of losing one of the sensory modalities. For example, losing vision not only alters the functionality of the visual cortex, but is also known to affect other sensory systems. Prior in-vitro studies of circuit-level changes in auditory cortex (ACX) showed global changes in synaptic transmission following light deprivation (Goel et al., 2006).

Methods
We investigated the functional consequences of these synaptic changes by examining sound-evoked responses of ACX neurons in-vivo using electrophysiological recordings and 2-photon imaging in mice that had been light-deprived for 5-10 days.

Results
We found fundamental alterations of key metrics of sound-evoked neuronal activity in dark-reared (DR) mice with observed differences between cortical layers. Driven firing rates, signal-to-noise ratios and slopes of rate-level functions of ACX neurons were significantly increased. Furthermore, there were significant alterations in frequency tuning properties of neurons within ACX.

Conclusion
We conclude that the functional alterations seen in the auditory cortex as a result of dark-rearing represents cross-modal compensation in the form of improved representation of auditory information. These changes may point towards potential for strategies such as visual deprivation that may be used for recovery of sensory function in other affected modalities, to possibly counter the effects of maladaptive changes in sensory processing.
Source Localization of the N1 Potential After Visual and Auditory Inhibition.
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Background
In response to an auditory stimulus train, the N1 potential of the
cortical auditory evoked potential in response to later stimuli of the train has been
shown to reduce in amplitude as compared to the first one. This phenomenon has
been related to lateral inhibition. The reduction in the amplitude of the N1 potential has
also been shown if an auditory stimulus is preceded by a stimulus in another sensory
modality, such as a visual stimulus. This indicates that the lateral inhibition may not be
specific to the auditory system, but originates from a source or multiples sources that
respond to multiple sensory modalities. This is of particular interest as the extent of
adaptation of the N1 potential has been shown to correlate with cochlear implant patient
speech perception ability. A non-auditory-specific source(s) of this adaptation may
dependently indicate non-auditory specific deficits in poor performing CI users. However,
these non-auditory-specific sources have not been identified yet.

Methods
Cortical auditory evoked potentials were recorded using a high-density
array of 160 electrodes in 10 normal hearing individuals. Stimuli for auditory-inhibited
responses were trains of five 1kHz tonebursts, with 1 second between tonebursts within
a train, and 12 seconds between trains. Stimuli for visual-inhibited responses were
a checkerboard pattern change, followed after 1 second by a 1kHz toneburst, with
12 seconds before the following visual stimulus. The topography of the auditory and
visual- and auditory-inhibited potentials were compared, as different cortical generators
produce different topographies at the scalp. Subject specific head model based current
source density algorithms were used to reconstruct cortical activity.

Results
Preliminary data suggest that the N1 potential of the response to the first
stimulus of a train of auditory stimuli (uninhibited response) has a different topography
to the response to later stimuli in that train (auditory inhibited response).

Conclusion
Further experiments are ongoing to verify our hypothesis.
Multi-Sensory Integration in Primary Auditory Cortex is Stimulus Timing Dependent and Alters Neural Synchrony
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Background
Central auditory neurons are influenced by activity-dependent inputs from non-auditory systems. In addition to acoustic stimulation, primary auditory cortex (A1) receives somatosensory input to areas previously considered to be exclusively "unimodal" (Hackett et al., J Comp Neurol, 2007). A1 neurons are capable of multisensory integration since spinal trigeminal (Sp5)-auditory stimulation modulates early event-related field potentials that are supra-additive (Lakatos et al., Neuron, 2007; Kayser et al., Neuron, 2005). The mechanisms underlying activity-dependent and spontaneous neural changes in A1 response properties following multi-modal stimulation remain largely unexplored. The present study examined the effects of combined auditory-Sp5 (bimodal) stimulation on tone-evoked, spontaneous firing rates and neural synchrony in A1 neurons in response to varied order and timing of bimodal stimuli.

Methods
Four-shank, 32-channel silicon electrodes were placed in guinea pig A1 to record tone-evoked responses and spontaneous unit activity before and after bimodal stimulation with alternating pairing order (tone-Sp5 or Sp5-tone) at various pairing intervals (40, 20, 10 and 0ms). Synchrony between neuron pairs was measured using cross-correlations (Vos et al., J Neurosci, 1999).

Results
Bimodal stimulation induced both facilitation and suppression of tone-evoked firing rates in A1. Greater levels of suppression were observed when Sp5 stimulation preceded tones by 40 and 20ms, while preceding Sp5 activation with tones robustly facilitated neural firing at all pairing intervals. Increased neural synchrony in A1 was observed when Sp5 preceded sound, but decreased when the pairing order was reversed (tone-Sp5).

Conclusion
The changes in responses to bimodal stimulation order observed here may reflect spike-timing dependent plasticity (Dahmen et al., J Neurosci, 2008). Thus, neuronal multisensory integration of A1 neurons may be influenced by temporal relationships of converging auditory and non-auditory sensory systems, which may be important in normal sensory processing and in central pathway aberrancy as observed in tinnitus.

This work was supported by NIH grants R03DC009893 (GJB) and P30DC05188.

Dependence of sensory responses on locomotion and vigilance along the central auditory pathway in awake-behaving mice
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Background
The dynamics of cortical network activity is influenced by the state of the animal as well as the dynamics of input coming from the sensory periphery, which is itself state-dependent. Much less is known about sensory responses in the cortex of awake-behaving preparations than in anesthetized or quietly immobile and not alert animals, particularly in the auditory system. In the visual cortex, locomotion (walking) causes a strong enhancement of sensory responses (Niell and Stryker, Neuron 65:472-9, 2010). However, it is not known how this effect relates to vision, cortical network dynamics, locomotion, or the state of vigilance of the animal.

Methods
To address these issues, we have developed a head-fixed awake-behaving preparation in which mice are free to walk on a custom made cylindrical treadmill based on Wienisch et al. (Springer protocols, 2012). We record the local field potential (LFP) and multiunit activity (MUA) across layers with a 16-channel laminar probe, or whole-cell membrane potential in auditory cortex while mice are engaged in passive behaviors or perform a Go/No-Go auditory discrimination task. We simultaneously record activity in the CA1 field of hippocampus using a separate 16-channel laminar probe and monitor the movements of the animal with an optical sensor and high-speed video.

Results
We find that, in contrast to the visual system, walking causes a significant suppression (~40%) of evoked responses to tones that is apparent in the onset, sustained, offset, and rebound suppression components. Also in contrast to the visual
system, where the locus of the locomotion effect is thought to be the cortex (Niell and Stryker, 2010), we find a similar suppression in the central nucleus of the inferior colliculus with or without bilateral silencing of the auditory cortices with muscimol. In the hippocampus, during walking we observe a strong and persistent theta rhythm (~8 Hz). During still periods we observe intermittent theta rhythmic activity as well as large amplitude irregular activity including intermittent fast ripples (~150 Hz) in the pyramidal layer often associated with sharp waves in other layers.

**Conclusion**
In ongoing experiments we are using this hippocampal rhythmic activity to define the state of vigilance of the animal, and exploring the dependence of sensory responses in auditory cortex on vigilance and behavioral task performance measures. In addition, we are further isolating the locus of the effect of locomotion on auditory responses with experiments in the auditory periphery.
Binaural sensitivity in human auditory cortex: analyzing the time course and spatial pattern of activity in fMRI
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Background
Although the representation of auditory space within the auditory cortex (AC) remains poorly understood, contemporary studies have combined psychophysical, neuroimaging, and physiological approaches in human and animal models to explore that representation and its dependence on behavioral and stimulus context. To that end, recent studies in our lab have used functional magnetic resonance imaging (fMRI) to study the dependence of human AC blood-oxygenation-level-dependent (BOLD) responses on the binaural characteristics of sounds, namely interaural differences of level (ILD) and time (ITD). The results demonstrate clear modulation of AC responses by ILD due to inhibition of the BOLD response by the ipsilateral ear, but surprisingly weak modulation of responses by ITD. Here, we examine whether the stimulus-specific time courses or multi-voxel activation patterns of AC BOLD response exhibit greater modulation by ITD.

Methods
BOLD responses were measured from continuously-acquired images (whole-head echo-planar imaging at 3T [Philips Achieva], TR=2s, 2.75x2.75x3 mm resolution) using an “event-related” paradigm that presented trains of narrowband or broadband clicks carrying ILD or ITD that varied parametrically across 1-s trials. Sounds were presented via insert earphones (Sensimetrics), and listeners tasked with detecting infrequent (~1/13 s) pitch-change targets. BOLD activity was analyzed on each individual’s cortical surface using FSL, FreeSurfer, and MATLAB. Three: separate analyses were conducted: univariate tests employing the general linear model (GLM), BOLD timecourse estimation, and multivoxel pattern similarity analyses.

Results
Consistent with previous results, response-ILD functions obtained from GLM analyses indicated significant reductions of the BOLD response for moderate ipsilateral ILD compared either to large contralateral or ipsilateral values. Response-ITD functions for both narrowband and broadband modulated sounds revealed little modulation of the overall BOLD response by that cue. Time-course analyses revealed significant response adaptation for both cues. Finally, multivoxel pattern analysis was used to assess the degree to which ILD and ITD sensitivity varied systematically at smaller spatial scales than detectable with standard univariate measures.

Conclusion
Together with past work, the results support a role for binaural suppression (as opposed to facilitation) in shaping the ILD sensitivity of AC bold responses. The effects of ITD, in contrast, are more subtle; they suggest that careful attention should be paid to the pattern of excitatory and inhibitory binaural processing in AC, and to potential modulatory effects of the behavioral task. [Supported by R01-DC011548, R03-DC009482]
Intracranial responses representing stop consonant voice onset time and place of articulation on the human posterior lateral superior temporal gyrus

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Background
Intracranial recordings performed in patients undergoing electrocorticographic (ECoG) evaluation for surgical remediation of medically intractable epilepsy have greatly enhanced our understanding of how speech is represented in the brain. Recently, we demonstrated using averaged trials that both voice onset time (VOT) and place of articulation (POA) of stop consonants were represented by high gamma ECoG activity recorded from posterior lateral superior temporal gyrus (PLST) (Steinschneider et al., Cereb Cortex, 2011, 21:2332-47). It is unclear from these analyses, however, whether representations of POA and VOT are reliable at the more environmentally relevant single trial level.

Methods
Here, we examined the reliability of high gamma ECoG activity to represent stop consonant POA and VOT at the single trial level using a sparse logistic regression classification analysis. The study was IRB approved and followed patient informed consent. Stimuli were synthesized syllables /ba/, /ga/, /da/, /pa/, /ka/, and /ta/. They were constructed to have identical first and fourth formant center frequencies and overlapping second and third formant transitions (e.g., /ba/ and /ga/ shared the same third formant transition while /da/ and /ga/ shared the same second formant transition). Differences in POA were mainly based on varying the spectral content of frication within the first 5 ms of the sounds. Spectra were maximal at 800 Hz for /ba/ and /pa/, 1600 Hz for /ga/ and /ka/, and 3000 Hz for /da/ and /ta/. VOT was 5 ms for /ba/, /ga/, /da/ (voiced stops) and 40 ms for /pa/, /ka/, and /ta/ (unvoiced stops).

Results
Classification analysis yielded highly reliable discrimination of VOT when comparing responses elicited by voiced and unvoiced stop consonants. In contrast, POA was not accurately represented in the ECoG when comparing the combined responses elicited by /ba/ and /pa/ versus those to /ga/ and /ka/, and /da/ and /ta/. There was also a poor and variable representation of POA when voiced and unvoiced stop consonants were analyzed separately. Additional correlational analysis indicated that responses to 800, 1600, and 3000 Hz pure tones could not reliably predict responses to syllables varying in their consonant POA.

Conclusion
VOT is a powerful cue for speech sound discrimination on PLST. In contrast, POA based primarily on spectral differences during frication are weak cues for neural discrimination in the same brain region. Discrimination of POA based on formant transition or combined cues may be more reliable. (KN and MS contributed equally to this work).
Neural entrainment to speech, a matter of time or frequency?
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Background
Spatially coherent neural activity measured by MEG, EEG, or LFP is entrained to the slow temporal modulations, i.e. envelope, of speech.

Methods
In the time domain, this entrainment is often decomposed into responses with different latencies, while in the frequency domain it is usually decomposed into neural oscillations in distinct frequency bands. Here, we compare time domain analysis and frequency domain analysis for the MEG responses to speech presented in a variety of auditory scenes.

Results
In the time domain, it is found that late (>~100 ms) auditory responses are robust to the acoustic background while early (<~50 ms) auditory responses are not. In the frequency domain, it is found that very low frequency activity (< 4 Hz, delta band) is robust to the acoustic background, while higher frequency activity (4-8 Hz, theta band) is not.

Conclusion
These results suggest that robust delta band activity reflects the audibility of speech while theta band activity reflects the intelligibility of speech.

Enhancement of the N1 Cortical Auditory Evoked Potential in Low-Level Background Noise
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Background
Though most people are familiar with the notion that background noise is detrimental to hearing, some reports indicate that low levels of background noise may actually enhance certain aspects of neural auditory stimulus encoding. These studies classify enhancement as an increase in the amplitude of the N1 cortical response in low levels of background noise relative to the N1 amplitude recorded in quiet. Because most cortical auditory evoked potential (CAEP) studies report an amplitude decrement with the addition of background noise, cases of enhancement are particularly intriguing and deserving of further investigation.

The present study sought to determine what conditions are required to elicit enhanced CAEPs in the presence of noise. Reports of N1 amplitude enhancement have thus far been confined to studies employing binaural listening conditions and relatively fast presentation rates. Therefore, we examined the effect of both signal presentation rate and differences in monaural vs. binaural mode of presentation.

Methods
CAEPs were recorded from sixteen young normally hearing listeners using a 64-channel electrode cap. The signal was a 450-ms naturally spoken /ba/ presented at a level of 80 dB with interstimulus intervals of 900 and 1900 ms. The signal was presented both in quiet and in the presence of a continuous speech-spectrum noise at +10 and at +30 dB signal-to-noise ratio (SNR).

Results
Binaural presentations using a fast stimulus rate elicited enhanced N1 amplitudes in background noise relative to the quiet condition. In contrast, slow presentation rates generally elicited decreasing N1 amplitudes with increasing background noise level. Thus, preliminary results suggest that both rapid presentation rates and binaural activity are crucial for the enhancement of N1 potentials in background noise. Further, the N1 enhancement was lost when low-frequency activity (i.e., <3 Hz) was filtered out, suggesting that the enhancement effect arises from low-frequency response activity.

Conclusion
Preliminary results indicate that background noise elicits enhanced N1 responses when stimuli are presented binaurally at a fast presentation rate. The enhancement appears to result from low-frequency activity in the CAEP response. The implications of the importance of monaural/binaural mode and rate of presentation on N1 CAEP enhancement as well as
the dependence of the enhancement effect on low frequency response activity will be discussed. [Supported by NIH-NIDCD (R03DC010914) and VA RR&D (GoEC4844C)]

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Multiplexed Single-Unit Codes for Sensory-Cognitive Mapping in Ferret Frontal Cortex During an Auditory Reversal Task
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Background
During auditory-guided behavior, flexible sensory-cognitive associations are needed for sudden changes in the behavioral meaning of sounds. The plasticity of neural coding in frontal cortex (FC) is likely to contribute to such behavioral flexibility. Single-units in FC exhibit spike-rate codes that separately represent target and distractor tones during a two-tone discrimination task (Fritz et al., 2010). However, it is not known if single-units in the ferret FC change their spike-rate codes for two discriminated tones when the behavioral meaning of the two tones are suddenly reversed.

Methods
Two ferrets were trained on a conditioned avoidance frequency discrimination task. Exactly the same sounds were presented in each block of trials, but the behavioral meaning of the sounds was reversed in alternating blocks. Tones from either a high- or low-frequency region were selected as the target for a given block of trials. The animal quickly learned to identify the target stimulus, and to stop licking a waterspout during targets to avoid a mild tail shock. Thus, behavioral responses to the task sounds were reversed for each consecutive block. After training, we recorded from single-units in the dorsal FC of the behaving, head-fixed ferret to study the plasticity of neural representations for high- and low-frequency sounds during the reversal task.

Results
We found neurons in dorsal FC that responded most strongly to the targets and more weakly to distractors, independent of the absolute frequency of the sounds. In other words, when the task was reversed, spike-rate codes for the task sounds also reversed, following the behavioral meaning rather than sound frequency. In some of the same cells, the target’s absolute frequency-range was tagged with characteristic spike-rate dynamics, allowing these neurons to encode both task meaning and absolute frequency of the stimulus.

Conclusion
Our results suggest that single-units in the dorsal FC of the ferret have multiplexed spike-rate codes representing orthogonal assignments of task meaning to sounds. This context-dependent plasticity may contribute to the behavioral flexibility required in a reversal task. We will also discuss our results on sensory (ie. tone frequency) vs. cognitive (ie. behavioral meaning) target-class coding in dorsal FC, the dependence of single-unit-based stimulus classification on the time-course of learning during task-reversal, and the coding of long-term memories to task-relevant sounds during non-task as well as task conditions.

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The neural mechanisms of the phonemic restoration in people with normal hearing
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Background
In our daily life, parts of speech are completely drowned out by noises; however, we can still hear the speech as if it has not been interrupted by noises. This is caused by the effect known for phonemic restoration. Impaired ability of phonemic restoration has been reported to be related to the difficulties in understanding speech in patients with hearing loss even when hearing level is corrected by hearing aids. However, the neural mechanism of the phonemic restoration is not clear. The aim of our study was to elucidate the neural mechanisms of the phonemic restoration in people with normal hearing.

Methods
Eight healthy male volunteers participated in our study. This study was approved by the Ethics Committee of Osaka City University, and all the participants gave written informed consent for participation. They were requested to carefully listen to and understand 3 stories with their eyes closed. The stories were constructed of recorded narratives in which several syllables were partly replaced by white noises for 300 ms. Although 2 of the stories were played forward (forward
condition), 1 story was played in reverse to assess the neural responses caused by listening to the noises (reverse condition). The neural responses while listening to these stories were measured by using MEG with high temporal resolution. The MEG data were analyzed using dual-state adaptive spatial filtering methods. The oscillatory power in the forward condition relative to that in the reverse condition was analyzed using statistical parametric mapping to make inferences at a population level.

**Results**
The MEG data from 1 participant were excluded from our analysis because the participant experienced hearing loss. We focused on the MEG data in the time range of 100 to 300 ms and in the frequency range of 3 to 5 Hz, since the sensor-level time-frequency maps showed event-related synchronization in this time-frequency area. The group analysis showed that the oscillatory power in the forward condition was higher than that in the reverse condition in 2 brain regions (\(P < 0.001\), uncorrected for multiple comparisons). One brain region extended from Brodmann area (BA) 32 and the other extended from BA6. The latter region seemed to overlap with Broca’s area.

**Conclusion**
We found two brain regions were related to phonemic restoration. Our findings may help clarify the neural mechanisms of the phonemic restoration as well as develop innovative methods for patients suffering from hearing loss.

**966 Interactions between areas in the forebrain of anaesthetized guinea pigs that elicit vocalizations**
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**Background**
Guinea pigs are gregarious animals with a well-characterized repertoire of 11 vocalizations. These are context dependent, communicating information about danger, identity and emotional state. In previous work guinea pig vocalizations were elicited by electrical stimulation of three brain areas, but the call types were not well classified. Here we investigate interactions between the areas that produce calls.

**Methods**
Five brain areas were electrically stimulated (60 Hz for 1.6 s) in anaesthetized animals: the anterior cingulate cortex, medio-dorsal thalamus, hypothalamus, amygdala and the midbrain periaqueductal grey substance. Spectral structure and pulse frequency of the evoked calls were analysed using SAS Lab Pro (Avisoft Bioacoustics).

**Results**
Eight distinct calls were classified. The structure of the calls was similar to those from spontaneously vocalizing animals. Five of the calls were produced only during electrical stimulation and three were produced after the end of the stimulation. Two of the post-stimulation calls were a series of “screams” gradually changing to “squeals” before fading away over a period of up to 30 s. These could be elicited from all five brain regions and showed a predictable response to changes in stimulation current. Concurrent bilateral stimulation of loci producing scream/squeal always had an additive effect, whereas stimulation of two loci giving during-stimulus calls was more complex. It appears that each post-stimulus vocalization area contributes to a shared system of excitatory re-entrant loops, that are involved in the generation of a call series.

**Conclusion**
Stimulation of the amygdala yielded two call types with contrasting emotional intent. To date, this has been reported only in primates. Our experimental protocol, therefore, provides a useful model for the investigation of the common mammalian vocalization pathways.
Alignment of Sound Localization Cues in the Nucleus of the Brachium of the Inferior Colliculus

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Background
Previous studies have demonstrated that although single neurons in the central nucleus of the inferior colliculus (ICC) are sensitive to multiple sound localization cues, they do not show a preference for spatially coherent cues. In the present study we measured the effects of misalignment of sound localization cues in the nucleus of the brachium of the IC (BIN) in unanesthetized marmoset monkeys. The BIN receives its predominant auditory input from the ICC and provides a major auditory projection to the superior colliculus, which contains a topographic map of auditory space.

Methods
Sound localization cues including interaural time differences (ITDs), interaural level differences (ILDs), and spectral shapes (SSs) were measured in each monkey. The results were used to impose aligned and misaligned combinations of cues on a set of broadband noise stimuli. The stimuli were presented to the same marmoset while recording spike responses in the BIN. We computed mutual information between the set of spike rates and stimuli containing either aligned or misaligned cues. In a subset of neurons noise ITD sensitivity and tone frequency response maps were also measured.

Results
The results can be summarized as follows: 1) A significantly larger number of BIN neurons encoded more information about auditory space when cues were aligned compared to misaligned. 2) This result holds in subsets of BIN neurons with or without ITD sensitivity. 3) In general tonal frequency tuning was broad (3-5 octaves). However, a subset of neurons that did not respond to tones below 2 kHz also had a preference for cue alignment.

Conclusion
The results suggest that ITD sensitivity is not necessary for cue alignment preferences in the BIN and posits an additional role for the interaction between ILD and SS cues. Alignment of sound localization cues in the BIN is hypothesized to play a role in the formation of the auditory space map found in the SC.
Spike-rate variance as the detection parameter for interaural incoherence
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Background
It has been shown that Inter-Aural Correlation (IAC) is linked to the variance of Inter-aural Time Difference (ITD) and Inter-aural Level Difference (ILD) and whilst normalised IAC can account for behavioural IAC discrimination performance so can models directly employing this variance.

The prevailing view, influenced by observations of binaural sluggishness, has been that this variance is too rapid for the binaural system to utilise. Attempts at identifying a neurometric correlate of IAC discrimination have consequently focused on changes in mean spike count, typically at the peak of ITD tuning curves.

We propose that IAC discrimination relies on variance spike rates on the slope of ITD tuning curves and compare computer simulations to midbrain neurons.

Methods
A physiologically-based hemispheric balance model was developed, where the fluctuations in the ratio between left and right brain activity served as the detection cue to a reduction in IAC from unity. Note that this ratio is stimulus power invariant (intrinsic normalisation). The analysis focused on neural populations with the same best ITD magnitude but opposite sign. Static changes in this ratio served as a cue to static changes in ITD. Comparisons were made with a model where changes in normalised mean spike rate at the ITD function peak served as the detection cue.

IC neural responses to binaurally presented noise stimuli of various IAC were recorded from anaesthetised Guinea pigs. Identical stimuli were presented twice, with ipsilateral and contralateral waveforms swapped so each neuron could serve as the model for a matching neuron with the same best ITD magnitude but in the opposite hemisphere aiding comparison to our models.

Results
Adjusting the model parameters to predict published behavioural ITD and IAC discrimination data, the hemispheric balance model required approximately 175 neurons in each hemisphere whilst two orders of magnitude more activity was needed to achieve the same performance with the model using mean group spike rate at the peak of their ITD functions. This adjustment also revealed a necessary neural time integration of 10ms, which is comparable with physiological estimates.

Conclusion
Hemispheric balance is invariant to stimulus power fluctuations but sensitive to interaural differences. Reduction in IAC leads to fluctuation in hemispheric balance and is a plausible IAC discrimination cue. Sensitivity to variance in ITD inherent in IAC stimuli supports recent evidence of a more rapid binaural neural system than suggested by “classic” behavioural binaural sluggishness.
In-Vivo Whole Cell Recordings Revealed Binaural Mechanism for the Facilitated Response in EI/F Neurons

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Background
Cells that receive excitation from one ear and inhibition from the other (EI cells) process interaural intensity disparities (IIDs), the cues for localizing high frequencies. EI cells in the inferior colliculus (IC) fire to contralateral stimulation, while ipsilateral stimulation at progressively higher intensities suppresses the spikes evoked by contralateral stimulation. Facilitated EI cells (EI/f) are a variation of EI cells in that facilitated spike-counts, more than those evoked by the contralateral signal, are evoked with binaural signals having low ipsilateral intensities.

Methods
To evaluate the binaural mechanism underlying the facilitated binaural response, we made whole cell patch-clamp recordings in 11 EI/f cells in the IC of awake Mexican free-tailed bats. Both spikes and postsynaptic potentials (PSPs) evoked by contralateral, ipsilateral and binaural signal were recorded. Moreover, excitatory and inhibitory synaptic conductances were derived from each response recorded in one cell.

Results
Based on both the PSPs and conductances, we propose the inputs that innervate these EI/f cells and show that the basic EI property was produced by innervation from a lower binaural nucleus, presumably the LSO. We then show that these cells also received an ipsilaterally evoked inhibition and an ipsilaterally evoked subthreshold excitation.

Conclusion
The facilitation resulted from two ways. 1) the ipsilaterally evoked excitation and a reduction in the ipsilaterally evoked inhibition due to contralateral stimulation. Thus, at small IIDs, with weak ipsilateral signals, the reduced inhibition allows the combined contralateral and ipsilateral excitatory inputs to generate a larger EPSP than would be evoked by the contralateral excitation alone, thereby generating the facilitated response. 2) only a reduction in the ipsilaterally evoked inhibition due to contralateral stimulation. This reduced inhibition produced a larger EPSP than that would be evoked by the contralateral excitation alone. Supported by NIH grant DC007856.

Establishing optogenetics in the auditory system of the Mongolian gerbil

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Background
The rodent Mongolian gerbil (Meriones unguiculatus) has been established as model organism for human hearing due to its similarly broad hearing range and the ability to localise low frequency sounds. However, by choosing the Mongolian gerbil as model organism we currently lack genetic tools and techniques already established in mouse or rat. Optogenetics is one example. Optogenetic techniques allow for specific targeting of individual brain nuclei and for control of electrical activity of neurons via light. Optical stimulation can circumvent off-target responses from passing fibres where electrical stimulation lacks precision.

Methods
Stereotactic injection of adeno-associated virus (AAVs) into auditory nuclei of anaesthetised gerbils was followed by an expression period of 2-3 weeks. Animals were sacrificed and brain slice preparations were performed. Whole-cell patch-clamp recordings from targeted auditory nuclei in acute brain slices were used to characterise kinetic parameters of the expressed channelrhodopsins. Immunohistochemistry and confocal scanning of paraformaldehyde-fixed brain slices served to measure the spread of the AAV-infection and to determine co-localisation of fluorescence-tagged channelrhodopsin with microtubule-associated protein 2 (MAP2).

Results
AAV mediated gene delivery reliably drove neuronal-specific expression of channelrhodopsins in the inferior colliculus (IC) and other auditory nuclei. Viral transduction of neurons in these nuclei with channelrhodopsin variants enabled us to control neuronal activity with light of appropriate wavelengths and characterise channelrhodopsins in the gerbil. Performing whole-cell patch-clamp recordings we determined maximum firing rates, firing reliability, and jitter of elicited
action potentials in current-clamp as well as transition from peak to stationary current (τin), current kinetics after light-off (τoff) and maximum photocurrents in voltage-clamp mode.

Conclusion
Our results are a first step to establish optogenetics in the auditory system of the Mongolian gerbil and might aid to decipher the intricate connectivity of auditory nuclei.

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Does Visual Stimulation Influence Spatial Tuning in the Central Nucleus of the Inferior Colliculus?
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Background
The central nucleus of the inferior colliculus (ICC) is one of the key stages in the auditory pathway. It is one of the main areas of convergence for both afferent and efferent fibers passing between the auditory brainstem and auditory cortex. It has many cells sensitive to particular auditory features, such as sound localization cues. It also receives indirect projections from multimodal structures, including the superior colliculus (SC) and the primary sensory cortices. This suggests that multimodal information is likely reaching the ICC. Further, since many ICC neurons are sensitive to sound direction cues, and vision is an inherently directional sense in most vertebrates, it would be natural for visual inputs to modulate responses to sound localization cues in the ICC. However, although its place in the auditory pathway is well-established, it has not been conclusively established that connections with other sensory modalities actually exist in the ICC.

Methods
The purpose of this study is to combine visual inputs with auditory inputs in order to look for any visual influences on ICC single-neuron responses in-vivo. To test for visual influences on ICC neurons responses from extracellular recordings in anesthetized Mongolian gerbils were analyzed.

Results
Paired directional visual inputs were combined with binaural sound inputs. A combinatorial analysis was carried out to look for synergistic, antagonistic, or non-selective visual influences on ITD and ILD response curves.

Conclusion
As the nexus of the auditory pathway, understanding the full range of influences on ICC neurons is critical to understanding how auditory information is processed and interpreted by the brain.

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Sensory responses to ethologically relevant auditory stimuli in the superior colliculus
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Background
Selective processing of stimulus elements in a complex environment lays the foundation for natural scene perception; however, the underlying neural mechanisms are not well understood. We have studied neural responses to natural acoustic stimuli in the midbrain superior colliculus (SC) of the echolocating bat, an animal that actively probes its environment by emitting ultrasonic vocalizations and listening to echo returns. Previous research on the bat SC has examined auditory evoked activity in superficial/intermediate layers (Valentine and Moss, J. Neurosci., 1997) and premotor activity in the intermediate/deep layers (Sinha and Moss, J. Neurosci, 2007). What remains elusive in the bat, and more generally for all mammals, is how sensory information is integrated into motor commands for perceptually guided behaviors. The purpose of the current study was to provide a broad framework for understanding sensorimotor processes in the mammalian SC. Echolocating bats serve as an excellent model for this line of study, because bats are auditory specialists that represent their 3D environment with sound and adapt vocal-motor behaviors in response to acoustic input.

Methods
In the current study, we examined SC encoding of dynamic echolocation signals employed by the bat while tracking moving prey items. Electrodes were lowered into the SC as the bat listened to natural sound stimuli. Responses were
later analyzed with respect to the depth of the recording site. By using this approach, it is possible to study how neurons in each layer of the bat SC respond to ethologically relevant auditory stimulation.

Results
Sensory responses to sonar vocalizations were elicited from SC neurons at superficial, intermediate and deep layers. Response latencies ranged from very short (<10 msecs) to very long (>45 msecs). We found selectivity in SC neurons to subsets of FM signals embedded in sound sequences over which call duration, interval, bandwidth and harmonic strength changed. Response selectivity was traced to correlated changes in acoustic features of sonar signals within the dynamic call sequences.

Conclusion
Sensory evoked responses were found across all depths of the bat SC, illuminating a process by which auditory information can affect ongoing motor planning in deeper layers. These results are novel in that they identify how auditory information is encoded in each of the layers of the SC, and they also provide a method to more broadly understand how sensory activity is represented in the SC of mammals.

Neural Coding of Interaural Level Differences with Bilateral Cochlear Implants
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Background
Interaural level differences (ILD) are by far the most potent sound localization cues presently available to bilateral cochlear implant users. Although the neural representation of ILD has been well-studied in normal hearing, virtually nothing is known about how the brain encodes ILDs of electrical stimuli.

Methods
We compared ILD sensitivity of single units in the inferior colliculus (IC) of barbiturate-urethane anesthetized cats and awake rabbits with bilateral cochlear implants. The cats were deafened one week before neural recordings, while recordings in rabbits lasted up to 6 months after deafening and implantation. Stimuli were trains of biphasic current pulses presented at rates <100 pulses/s. Current levels were systematically varied in each ear starting just below threshold and spanning the dynamic range, yielding a matrix of firing rate as a function of stimulus level in each ear. Rate-ILD curves at different mean binaural levels (MBL) were extracted from each matrix.

Results
Neural ILD sensitivity evoked by electric stimulation was qualitatively similar to that previously observed in normal hearing animals. Stimulation of the contralateral ear evoked excitation in the vast majority of neurons. The predominant effect of ipsilateral stimulation could be excitatory (“EE” response), inhibitory (“EI”), or weak (“EO”). A few neurons responded nonmonotonically with increasing level to contralateral and/or ipsilateral stimulation. EI neurons showed a monotonic dependence of firing rate on ILD that was relatively invariant with changes in MBL, thereby providing a robust code for ILD. In contrast, rate-ILD curves for EE neurons were nonmonotonic and highly dependent on MBL, providing an ambiguous code for ILD that degraded at high MBLs. Such neurons might still provide a distributed code for ILD if their ipsilateral and contralateral thresholds varied systematically across the neuron population.

The prevalence of the various binaural response types appeared to differ between the two preparations. EE was most common in cat, followed by EO and EI. Based on a small sample, the EO type predominated in rabbit, followed by EI and EE.

Conclusion
Our results suggest that the predominant code for ILD may differ in acutely-deafened, anesthetized cats vs. chronically-implanted, awake rabbits. Several factors may contribute to this difference, including effects of anesthesia, species differences, or sampling biases. Additionally, in the chronic rabbit preparation, the balance of excitation and inhibition may be influenced by prolonged deprivation of acoustic input and by electric stimulation during daily recording sessions.

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Investigating the Effect of Microphone Position on Sound Localization in Bilateral Cochlear Implant Users

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Background

Bilateral cochlear implants (BiCIs) have become a standard of care in many clinics, in an effort to improve spatial hearing skills in profoundly deaf individuals. However, BiCI users demonstrate poorer sound localization accuracy compared to normal hearing (NH) listeners. One of the many factors that might contribute to poorer performance is the behind-the-ear microphone placement of these devices. Such placement does not maximize the ability to capture the natural filtering properties of the head/pinna known to be important cues for sound localization. Here, we evaluated the effect of microphone placement on sound localization performance in BiCI users. We hypothesized that the most natural position, in-the-ear placement of microphones, will capture spatial cues that are unavailable in standard CI fittings, improving performance.

Methods

Eight postlingually deafened BiCI users participated in this study. Virtual acoustic space (VAS) techniques were used to capture individualized acoustic transfer functions (ATFs) recorded for 19 loudspeaker locations, spaced 10° apart along the horizontal plane. Microphones were positioned to reflect those currently used by clinics: (1) in-the-ear (ITE); (2) behind-the-ear (BTE); or (3) on the shoulders (SHD). ATFs were used to generate VAS stimuli consisting of a pink noise train (four burst each 170 ms). Stimuli were delivered to the CI user’s speech processors through direct connect cables. A sound localization task was conducted to evaluate performance for each microphone position condition. Free-field localization using the subject’s clinical processors was also tested.

Results

Subjects were able to localize all VAS stimuli, except one who indicated that the stimuli did not sound externalized. Results demonstrate that placing the microphone in-the-ear resulted in improved localization accuracy compared to the free-field and other microphone conditions. Accuracy and sensitivity decreased substantially in the SHD condition. In all conditions, localization performance was still less accurate compared to NH listeners.

Conclusion

We found ITE microphone placement provides a small improvement in sound localization over BTE. However, the results also indicated that ITE placement is not enough to restore localization to NH performance. Additional factors need to be considered for improvement of sound localization in BiCI users, such as the restoration of binaural timing cues known to be important for accurate sound localization. Combining the preservation of acoustic temporal fine structure with bilateral synchronization between processors may provide these cues and improve BiCI sound localization. Supported by NIH-NIDCD (R01003083) and (P30 HD03352).

The Effect of Interaural Frequency Mismatch on Correlation Change Detection in Bilateral Cochlear-Implant Users

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Background

The insertion depth of a cochlear implant (CI) may differ across the ears in the case of bilateral implantation. This will cause an interaural frequency mismatch of electrical stimulation that may affect binaural processing, which is thought to require matched frequency inputs. While previous studies have investigated the effects of interaural frequency mismatch on the perception of static interaural differences using constant-amplitude pulse trains, we extended that work to investigate interaural correlation change detection in modulated stimuli.

Methods

In experiment 1, correlation change discrimination just-noticeable differences (JNDs) were measured in six CI listeners. The stimuli began as a 50-Hz narrowband noise. The envelope was extracted, biphasic monopolar electrical pulses sampled the envelope at 1000 pulses per second (pps), and amplitudes were compressed as typically occurs in speech processing strategies. Starting from a pitch-matched pair of electrodes, interaural frequency mismatches of \( \Delta = 0, 2, 4, \) and 8 electrodes were tested. JNDs were measured for correlated and uncorrelated reference stimuli. In experiment 2, the percentage of correct discriminations between correlated and anti-correlated (i.e., half-period phase shifted) stimuli was measured as a function of pulse rate. In experiment 3, a fusion experiment was performed to measure the subjective
diffuseness of the correlated, uncorrelated, and anti-correlated stimuli. Parallel simulation experiments using acoustic pulse trains were performed in normal-hearing listeners.

Results
In experiment 1, all listeners were sensitive to changes in correlation from correlated reference stimuli for $\Delta = 0$. JNDs were relatively worse for changes from uncorrelated references. Increasing $\Delta$ caused increased JNDs; however, most listeners could tolerate a shift of four electrodes (3 mm) before JNDs increased. In experiment 2, listeners could almost perfectly discriminate correlated and anti-correlated pulse trains at 100 pps. However, performance dropped to chance by 500 pps. In experiment 3, uncorrelated and anti-correlated stimuli were perceived to be more diffuse than correlated stimuli.

Conclusion
An interaural frequency mismatch of more than 3 mm is detrimental to processing of changes in interaural correlation, consistent with previous mismatch studies using monopolar stimulation. Given that listeners were not sensitive to changes in phase of the electrical stimuli (correlated vs anti-correlated) at rates typically used in speech processing strategies, this demonstrates a significant hurdle in presenting binaurally-relevant fine-structure information to CI users.

Lateralization of Interaural Level Differences in Multi-Channel Electrical Stimulation
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Background
Improved sound localization is one of the benefits associated with use of bilateral cochlear implants (CIs). Modern speech processors cannot reliably encode all of the localization cues available to normal-hearing listeners. Therefore, CI listeners rely mostly on interaural level differences (ILDs) to produce a spatial impression of the surrounding environment. The thresholds (T) and the "most-comfortable" (C) levels obtained during the mapping procedures often differ across the electrodes within the ear and across the ears, resulting in a different spread of excitation at different cochlear locations. Moreover, real world sounds can activate multiple electrodes simultaneously. It is unclear if multi-electrode stimulation with current spread can produce a centered and punctate auditory image in a CI user's head, which could be used as a baseline to create a consistent spatial map across frequencies.

Methods
We investigated the perceived intracranial location of direct electrical stimulation in a group of bilateral CI listeners. For five electrodes in each ear, T and C levels were obtained. Stimuli were monopolar, 1000 pulse/sec pulse trains that were 500 ms in duration. They were presented bilaterally at 70-80% of the dynamic range using single-electrode pairs and several multi-electrode combinations, including three widely-spaced pairs, three closely-spaced pairs, and five-electrode pairs. Listeners reported the position of the perceived intracranial image at ILDs of 0, ±2, ±5, and ±10 current units (cus). Lateralization ability with inconsistent ILDs across frequencies was studied in normal-hearing listeners to simulate lateralization with multi-electrode stimulation in CI listeners.

Results
The shape of the lateralization curves was highly variable for single- and multi-electrode stimulation, both within and across listeners. This often resulted in an offset in the perception of a centered image for 0-cu ILD. Adjusting the levels of individual electrode-pairs within a multi-electrode combination with the offsets measured in the single-electrode pairs shifted the location of the perceived sound at 0-cu ILD. This shift resulted in a more centered image for 7/10 multi-electrode combinations in 4 listeners.

Conclusion
The results suggest that auditory spatial maps are distorted for single- and multi-electrode stimulation in bilateral CI users. The perception of a centered image for 0-cu ILD can be improved by adjusting the levels of individual electrodes within a multi-electrode combination based on the single-electrode data. These data have implications for bilateral clinical mapping strategies.
Place Coding In The Cochlear Apex
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Background
The majority of cochlear implant psychophysics has been conducted with users of electrodes that are typically inserted approximately 1.25 turns into the cochlea. From work with these devices, it is commonly accepted that a change in place of stimulation corresponds to a change pitch.

Med-El produces two long (31.5 mm) electrode arrays (the Standard and FLEXSoft), which are inserted deeper into the cochlea than traditional electrode arrays, providing an opportunity to study the properties of the cochlear apex. Pitch scaling studies with the long Med-El arrays show that some patients describe the most apical electrodes as having a similar pitch while others report that pitch lowers continuously across the array. One explanation is that as the electrodes approach the apex, more peripheral processes are stimulated and therefore the reported pitch change may not be of the same quality as place changes elsewhere in the cochlea (similarly to how place and rate changes do not produce the same quality of pitch change). Therefore, we conducted an investigation into the perceptual quality of a change in place of stimulation across 31.5 mm electrode arrays using Multi-Dimensional Scaling.

Methods
Stimuli consisted of equally loud single electrode, 5000 pulse-per-second pulse trains. Nine stimuli represented electrodes numbered 1-9. Note, in the Med-El system, 1 is the most apical electrode out of 12. In a given trial, a pair of stimuli was presented to the listener who had to scale how different the two sounds were. All pairs of stimuli were compared 10 times. Nine subjects were tested.

Results
An INDSCAL analysis shows that in two dimensions, each electrode is plotted in order in a horseshoe configuration, suggesting that even with a deep insertion, a change in place yields a change in place pitch throughout the array. However, the perceptual distances between the most apical electrodes are shorter than the perceptual distances between other electrodes. Analysis of individual subjects ALCSAL plots suggest that most patients results are typical of the INDSCAL analysis; however a number of patients show a perceptual clustering near the apex.

Conclusion
Deep insertions provide continuous place pitch throughout the cochlea. From a place pitch perspective, the extended length of the Med-El Standard and FLEXSoft array provides additional benefit relative to shorter electrode arrays. However, the ideal electrode configuration might involve greater spacing between electrodes in the apex than the rest of the array.

Ipsilateral and contralateral masking in bilateral cochlear implant users
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Background
It has been proposed that cochlear implant (CI) users’ spectral resolution can be improved by distributing adjacent spectral channels across ears, an approach referred to here as interleaved processors. However, the benefit of interleaved processors depends strongly on selecting truly independent channels within and across ears. Ipsilateral and contralateral masking may be used to measure the spread of excitation (SOE), which can be used to identify independent spectral channels. For normal-hearing (NH) listeners, peak ipsilateral and contralateral masking occurs when the probe and the masker are approximately the same frequency, and the spread of masking is narrower with contralateral than with ipsilateral masking. Two factors may cause CI listeners’ contralateral masking functions to differ from those of NH listeners. First, stimulating matched cochleotopic locations in each ear may not yield the same perceived pitch. Second, because CIs directly stimulate the auditory nerve, much of the cochlear mechanics that may be responsible for the greater spread of ipsilateral masking functions are circumvented. The goals of this study were to determine whether the peak of contralateral masking for CI listeners occurs when the probe and masker yield the same pitch percept and whether the SOE for CI listeners was reduced with contralateral maskers.

Methods
Pitch matching was performed by stimulating a reference stimulus in one ear and adjusting the stimulation site of an equally loud probe in the opposite ear until the pitch matched across ears. Contralateral and ipsilateral masking was
measured using a modified Bekesy tracking procedure; a 20 ms probe was temporally embedded in a 500 ms masker. Maskers were presented at the most comfortable loudness level and loudness matched across ears. The ipsilateral masker was placed at the peak of the contralateral masking function.

Results
Preliminary data suggest that contralateral and ipsilateral masking functions differ considerably across patients and across stimulation sites. Despite that variability, the results indicated that the peak of contralateral masking occurred at approximately the same location as the perceived pitch matches. The results also indicated that, despite circumventing much of the cochlear mechanics, SOE was narrower with contralateral than with ipsilateral masking.

Conclusion
The relatively narrower contralateral SOE suggests that the number of independent channels can be increased by moving some stimulation sites to the opposite ear. The results also suggest that, whether using interleaved processors or standard clinical processors, pitch matching can provide an efficient method to optimally align stimulation across arrays.

Use of a dual-task paradigm to assess listening effort in a CI simulation
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Background
Many bilateral or bimodal users of cochlear implants (CIs) report that it is easier to listen when both devices are active than with only a single device, especially in noise. However, there is currently no method to quantify listening effort in these patients. Here we used a CI simulation to test the feasibility of using a dual-task paradigm to measure listening effort. In a dual task paradigm, the listener divides attention between a primary and secondary task. As the primary task becomes more difficult, fewer cognitive resources are available for the secondary task, generally resulting in poorer performance. Our hypothesis was that when listeners perform a sentence-recognition task, as signal-to-noise ratio (SNR) decreases, performance on a secondary task will decrease due to an increase in listening effort.

Methods
Normal-hearing young adults participated. The primary task was to repeat sentences processed through an 8-channel CI simulation. AzBio Sentence Lists were presented in quiet and noise (+16, +12, and +8 dB SNR). The secondary task was to recall 7 digits presented visually before each set of 4 sentences. As a control, both the sentence-recognition and digit-recall tasks were tested singly. The NASA Task Load Indices for Effort, Mental Demand, and Performance were completed in each test condition.

Results
On the primary sentence-recognition task, performance did not differ between the single- and dual-task paradigms. Sentence-recognition scores decreased significantly as the SNR deteriorated. In contrast, the digit-recall scores were significantly lower in the dual-task paradigm, suggesting an increase in listening effort. However, no significant changes occurred in digit-recall as the SNR deteriorated. Finally, the NASA ratings indicated a significant perceived increase in listening effort and mental demand and decrease in performance as the SNR deteriorated.

Conclusion
A dual-task paradigm may be useful for quantifying listening effort because, when compared to the single-task control 1) performance on the primary sentence-recognition task was not affected, but 2) performance on the secondary digit-recall task decreased significantly. However, scores on the digit-recall task did not decrease as the SNR deteriorated. This may be due to the test parameters utilized here, or because the digit-recall task was suboptimal for observing smaller changes in listening effort, particularly within individual listeners. We are modifying our protocol to better understand these issues, and are testing the efficacy of the dual-task paradigm in CI recipients.
Self-selected and pitch-matched frequency tables in users of bilateral cochlear implants
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Background
Bilateral cochlear implants (CIs) provide many benefits including improved speech-understanding abilities in noise and sound-localization. Currently, bilateral CIs are fit by programming each ear independently, with minor adjustments to balance loudness between ears. While this procedure yields bilateral benefit in most patients, it cannot account for between-ear mismatches in insertion depth or neural survival. To address this issue, we have been exploring the use of real-time manipulations of the frequency table as a possible tool to aid the fitting of bilateral cochlear implants. Our previous data suggest that a subset of individuals select a frequency table in one ear that differs from the standard table worn in the contralateral ear. Our goal was to determine whether such selections are made because the listener is attempting to match the pitch percepts heard in each ear.

Methods
Participants included bilateral recipients of the Advanced Bionics (AB) device. All participants were initially presented speech via loudspeaker, and were instructed to adjust the frequency table until speech intelligibility was maximized. These selections were made both unilaterally, and with the contralateral CI active. All frequency-table selections were made in real time using custom software. Then, for each electrode in the array, a pure tone was presented via loudspeaker at the center frequency of the standard frequency table for one ear. In the contralateral ear, patients adjusted the frequency table until the pitch percepts were matched. These selections were then combined into a global pitch-matched table.

Results
Within a given listener, any shifts in the frequency table to bilaterally match the pitch percepts were largely consistent in the apical, mid, and basal end of the array. Taken together, the global pitch-matched table often differed from the table selected to maximize speech intelligibility. Generally, these pitch-matched tables were shifted higher in frequency than the self-selected table, which was usually shifted higher in frequency than the standard.

Conclusion
When listeners select a preferred frequency table that differs from the standard, they may be trying to select a signal that is as natural-sounding as possible without hindering speech intelligibility. Here, the pitch-matched tables rarely lined up perfectly with the self-selected tables for speech stimuli, especially in those cases where the self-selected speech table differed from the standard. We are currently investigating how performance on tests of speech intelligibility and sound-localization are affected by use of these frequency tables.
Absent cortical ITD representation in congenital single-sided deafness
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Background
Despite substantial reduction of ITD sensitivity in the cortex of congenitally deaf animals, cells sensitive to ITD changes were still observed (Tillein et al., 2010, Cereb Cortex 20:492-506). The auditory cortex of subjects with early single-sided deafness receives asymmetrical input leading to extensive reorganization of cortical aural dominance with aural preference for the hearing ear (Kral et al. 2012, Brain, in press). The recent study focuses on cortical ITD processing in this asymmetric hearing condition.

Methods
Two adult cats with congenitally unilateral deafness were compared to four adult deaf and four adult hearing control cats. Controls were acutely deafened by intracochlear application of neomycin. All animals were acutely stimulated with charge-balanced biphasic pulses (200µs/phase) in wide bipolar configuration through a custom made cochlear implant inserted into the scala tympani on either side. Cortical regions which were most responsive as defined by local field potential mapping were subjected to recordings of multi-unit activity using 16 channel electrode arrays (Neuronexus probes). Sensitivity to ITDs between -600 and +600 µs was tested with pulse trains (500 Hz, 3 pulses) at intensities of 0 – 10 dB above brainstem response thresholds. Template ITD functions (Tillein et al., 2010) were fitted to the data and the templates were statistically compared. ITD functions which did not fit significantly to the templates or had a modulation depth below 50 % were categorized as unclassified.

Results
Results demonstrate a significant increase in the number of unclassified but responsive cells with respect to deaf and hearing animals. These were found in more than 50% of all recording sites. However, when compared to deaf animals a significant increase in responsive sites was observed, so that with respect to responsiveness single-sided animals compared well to normal hearing controls. In consequence, cortical cells remained responsive to binaural stimulation, although the response was not ITD specific. Modulation depth of ITD functions in classified responses was additionally significantly smaller than in deaf and in hearing animals.

Conclusion
This study demonstrates that monaural deprivation leads to a distinct decrease of sensitivity to variation of binaural cues, whereas general responsiveness to binaural stimulation is comparable to hearing animals. This finding is in strong contrast to the situation in binaural deafness and demonstrates the detrimental influence of unilateral deafness on binaural feature sensitivity and consequently on the localization ability.

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COMPARISON OF SIGNAL-DETECTION AND GAP-DETECTION THRESHOLDS WITH FOCUSED COCHLEAR IMPLANT STIMULATION
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Background
With focused stimulation, such as with the tripolar electrode configuration, detection thresholds often vary substantially across electrodes for a given listener. In addition, we have previously shown that channels with high thresholds exhibit poorer frequency selectivity than those with low thresholds. These two findings could reflect variations in the health of auditory nerve fibers across the neural array, or, alternatively, variations in electrode distance from the modiolus. Differentiating the roles of electrode distance to nearby neurons and neural health (viability) is difficult for two reasons: 1) modern technology cannot image spiral ganglion neurons, and 2) imaging of the location of the electrode array is costly. Here we describe the measurement of gap detection thresholds, with the rationale that they might be elevated by the altered temporal discharge patterns associated with neural degeneration, but are less likely to be affected by the distance of the electrode from the modiolus.

Methods
Behavioral thresholds with focused stimulation were obtained for seven adult CI users to identify two channels with low and two channels with high thresholds for further testing. Gap detection threshold was then measured on the four test channels with the focused partial tripolar electrode configuration (sigma = 0.75). A 2-interval forced choice, 2-down, 1-up
adaptive procedure was used. Stimuli were 194 us/phase, 1031 pulses-per-second pulse trains lasting 200 ms. Both the train duration and level were roved. Gap detection thresholds were obtained at the most comfortable listening level for each channel unless that level could not be reached, in which case stimuli were presented at a medium-soft level that was loudness balanced across test channels. Thresholds were measured four or five times for each stimulus condition.

Results
The results showed a trend for channels with high thresholds to be basally located and to have longer gap detection thresholds, but there were clear exceptions to this relationship.

Conclusion
It is suggested that gap detection thresholds may help differentiate the effects of neural health and modiolar position on tripolar detection thresholds.

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Assessing Spatial Channel Interactions in Cochlear Implants
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Background
The main drawback of modern multi-channel cochlear implants (CI) is that the spatial selectivity of the excitation pattern is severely limited by the widespread nature of electric fields in the extracellular tissue. Modern CIs have the flexibility to electrically stimulate the auditory nerve and to capture various electrical signals inside the cochlear lumen.

Methods
This study examines spatial channel interactions at the electrical and neural level in a group of Advanced Bionics HiRes90K users. All subjects were implanted with the Helix array. The EFI measurements were made with the EFIM research tool. EFI data were obtained for all electrodes. ECAP recordings were done using the RSPOM software. SOEs were obtained from 4 different electrodes (3-7-11-15) using a variable recording location paradigm. SOEs were collected with 4 different loudness levels.

Results
Width, slope and symmetry of spatial excitation patterns derived from these physical (EFI) and physiological (SOE) measures will be compared.

Conclusion
EFI recordings allow a finer interpretation of neural SOEs as it teases out the effects of electrical spread within the cochlea. Taken as a whole, this study serves as a direct comparison and identification of inter- and intra-subject variability in spatial profiles.
Current Focusing Improves Speech in Noise and Spectral Resolution in Cochlear Implant Users

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Background

Although each electrode in a cochlear implant provides independent information, the spreads of excitation from adjacent electrodes overlap such that multiple electrodes stimulate the same neural populations. To reduce channel interaction and spread of excitation, current focusing techniques have been proposed in which multiple electrodes provide simultaneous stimulation out of phase to restrict current spread. One current focusing technique, partial tripolar stimulation (PTP) involves stimulating on two flanking electrodes in the opposite phase of a central electrode. Previous work has shown that the spread of excitation from PTP stimulation can be narrower than monopolar (MP) stimulation. In the present study, we implemented MP and PTP stimulation in novel speech processing strategies. Patients were tested on speech in noise and spectral resolution tasks to evaluate differences between the strategies.

Methods

Two novel sound coding strategies were implemented into an Advanced Bionics body worn speech processor. Each strategy consisted of 14-channel CIS to electrodes 2-15. The two strategies were identical in all parameters except for stimulation mode (MP stimulation for Exp-MP strategy and PTP for Exp-PTP strategy). M-levels for each electrode were carefully loudness balanced across the two strategies to ensure an equivalent spectral shape. Performance with each strategy was evaluated with HINT sentences, digits in noise and a new test to measure spectral resolution, the Spectral-temporally Modulated Ripple Test (SMRT). The SMRT consists of a spectral ripple whose phase drifts in time, eliminating potential loudness or edge effects of traditional spectral ripple tasks. Furthermore, the SMRT prevents the listener from performing the task by attending to a single electrode. Patients were evaluated on their spectral resolution with the SMRT test using both Exp-MP and Exp-PTP strategies.

Results

The average improvement in Speech Reception Thresholds (SRTs) with Exp-TP was 2.7 dB for HINT sentences in noise and 3 dB for digits in noise. All subjects performed better with Exp-TP with HINT sentences. One subject performed similarly with both strategies for the digits in noise task while all others performed better with the Exp-TP strategy. In pilot data, all subjects tested have shown better SMRT scores (i.e. better spectral resolution) with the Exp-TP strategy.

Conclusion

Current focusing (specifically PTP stimulation) provides better speech understanding in noise and better spectral resolution. Although further optimization of the strategies are needed, current focusing is a promising tool for better performance in future cochlear implant strategies.

Stimulation of the cochlear apex with partial-bipolar asymmetric pulses: Effects on speech intelligibility and pitch contour identification by cochlear implant listeners

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Background

Bipolar asymmetric pulses presented on the most apical channel of a cochlear implant (CI) can extend the range of both place and temporal pitch cues available to CI listeners (Macherey et al., JARO, 2011). Here, we investigate whether the implementation of such pulses may improve the perception of speech and the ability to identify pitch contours of CI subjects.

Methods

Ten users of the Advanced Bionics CII/HiRes 90k device took part in two sets of experiments. In the first, we measured consonant identification in a VCv context and sentence recognition using low-pass filtered IHR sentences for two speech-processing strategies. The two strategies were identical in all respects except that the most apical channel delivered either monopolar symmetric or partial-bipolar asymmetric pulses. In the second set, subjects had to identify a five-note pitch contour in a five-alternative forced choice task. The spacing between notes was three semi-tones and several ranges of fundamental frequencies were tested. Performance was measured in three conditions: (1) for a subset of channels delivering high-rate monopolar pulse trains sinusoidally-amplitude modulated (SAM) at the fundamental frequency (F0); (2) for a partial-bipolar pulse train presented at the apex at a rate equal to F0; and (3) for a combination of
(1) and (2). An additional condition presented pulses at the F0 rate to the apex, but, unlike condition (2), used monopolar stimulation.

Results
The results of the speech tests showed no difference in performance between the two strategies. The pitch contour test showed that a partial bipolar asymmetric pulse train presented at F0 on the apical channel can improve the identification of pitch contours, even when combined with monopolar SAM pulse trains on other channels. However, replacing the partial-bipolar asymmetric pulses by monopolar symmetric pulses on the apical channel yielded similar benefits.

Conclusion
The results of this acute study revealed no advantage of having partial bipolar asymmetric pulses instead of monopolar symmetric pulses on the apical channel of a CI. They also suggest that temporal pitch perception may be improved by an explicit representation of F0 delivered to a single channel (which in the present study was at the apex).


Refractory Properties of Cochlear Implant-Induced Spiking in Auditory Nerve Fibers are Dependent on Location of Stimulation and Voltage-Gated Channel Type Distribution
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Background
Experimental work has demonstrated that auditory nerve fibers (ANFs) of cats cannot fully respond to high rates of electrical stimulation, thus reducing the information transfer to the brain. Miller et al. (JARO 2001) have shown that a limiting factor of the reduced spike information transfer can be attributed to the neuron's refractory period. A computational model of a node of Ranvier of the ANF (Negm and Bruce, EMBC 2008) suggested that low-threshold potassium (KLT) and hyperpolarization-activated cyclic nucleotide-gated cation (HCN) channels (Yi et al., J Neurophysiol 2010) might be responsible for a larger refractory period (Negm and Bruce, in prep.).

Methods
We extend that work with a simulation study taking into account ANF morphology (Woo et al., JARO 2010) to consider the differential spiking activity as a function of 1) the location of electrical stimulation and 2) nodal channel composition at important locations along the ANF. Specifically, we test three ANF models variants: A) only fast Nav and delayed-rectifier Kv at all nodes, B) with the addition of KLT & HCN channels (Yi et al., J Neurophysiol 2010) at the first peripheral node and on the nodes of Ranvier neighboring the soma and C) by expanding the distribution of KLT channels to all nodes (Bortone et al., Hearing Res 2006).

Results
In general, we observed the absolute refractory period of model C to be the greatest followed by model B, then by model A. Models B and C contrasted with model A by having a greater probability of spike initiation at the location of stimulation. Model A did not show a strong relative refractory period at its peripheral nodes. We argue that the washout of the relative refractory period in this region was dependent on the low correlation between the location of the stimulating electrode and the location of spike initiation.

Conclusion
Preliminary results indicate that model C is most consistent with the published physiological data. In addition to the KLT & HCN channels of model C, other ion channel types may be necessary to explain all aspects of refractory behavior observed in vivo.

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Qualitative ratings of current focusing by cochlear implant users
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Background
Channel interaction is believed to be a limiting factor in the number of effective spectral channels available to cochlear implant patients. Current focusing (determined by coefficient $\sigma$) is expected to reduce channel interactions, which could lead to better speech performance under difficult listening conditions. In a previous study (Landsberger et al., 2012) we related the reduction in spread of excitation from current focusing on one electrode with qualitative ratings of the tone perceived by CI users. We found a relationship between the reduction of spread of excitation and qualitative ratings. These results were exciting in that an adjective scaling task could be used as a quick (and possibly clinically implementable) test to predict who would benefit from current focusing. However, in the previous study, only one location was tested in each of six patients. In this follow-up study, we measured qualitative ratings at many locations across the array on multiple subjects to determine the variability of qualitative ratings across the array within subjects. Furthermore, we collected forward masked spread-of-excitation measures at a number of locations to verify if the relationship between spread of excitation and qualitative rating is maintained with more data points. If this is the case, then qualitative ratings could be an easier and faster way to make these measurements reliably.

Methods
Seven users of the Advanced Bionics CII and HiRes 90k cochlear implants participated in this study. Spread of excitation for electrodes 4, 6, 10 and 12 was measured, using a 2IFC forward masking task. Qualitative ratings were measured for electrodes 2, 4, 6, 8, 9, 10, 12 and 14 using only two $\sigma$ values (0 and 0.75). The same adjectives as in the previous study were used for ratings (clean/dirty, pure/noisy, high/low, full/thin, flute/kazoo).

Results
Preliminary results suggest that spread of excitation is related to the qualitative ratings used in most electrodes. As in the previous study, the Clean / Dirty Index, seems to be the most closely related to the reduction of spread of excitation.

Conclusion
It seems possible to quickly predict spread of excitation by using qualitative ratings. Of all of the tested adjective pairs, the clean/dirty adjective pair best predict the amount of spread of excitation in that given electrode.

Musical Instrument Identification in Cochlear Implant Users: A Psychophysical Study of Acoustic Changes
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Background
Cochlear implants (CIs) offer many deaf individuals greater opportunity to interact with their environment, but there are still shortcomings in the user’s ability to perceive sound, particularly music. CI users perceive rhythm, but struggle
distinguishing between instruments. This study examined CI users’ ability to hear musical changes and explored the viability of using standard clinical equipment to measure auditory evoked potentials (AEP) in response to instrument changes.

Methods
Two same-different tests were designed to examine CI users’ discrimination of instrument stimuli. Sample notes from 5 instruments (clarinet, flute, violin, saxophone, and trumpet) were trimmed to only include the center 500ms. The first test concatenated random combinations of these instrument samples and asked subjects to identify the sounds as same or different. The second test spaced the combinations by 500ms and again asked users to identify whether there were differences between sounds. Ten experienced adult CI users and four adult normal hearing (NH) subjects were administered both tests. Percent correct scores and sensitivity indices (D’) were compared within and between groups. AEPs were measured on 4 NH subjects in response to clarinet-trumpet changes using clinical ABR equipment and a simple 3-4 electrode montage.

Results
CI subjects had lower discrimination scores on all tests and instrument combinations. However, both groups followed similar patterns in terms of which instruments seemed to be the easiest or most difficult to differentiate: clarinet-saxophone combinations were most difficult (NH D’=1.70, CI D’=.85), while both groups correctly identified flute-trumpet combinations over 99% of the time. Normal hearing D’ scores for the spaced tests were .44 standard deviations lower than concatenate scores (p=.03), while CI D’ scores were equal across the two tests. Statistically significant differences between NH and CI D’ scores were observed for trumpet-violin, flute-saxophone, and clarinet-saxophone combinations. AEPs were successfully recorded in four NH individuals in response to clarinet-trumpet changes. The stimulus onset responses and acoustic change complexes were observed.

Conclusion
CI subjects have a baseline lower capacity than NH for instrument discrimination; however, both groups struggle with the same instrument combinations. Several CI subjects who scored poorly on instrument identification tasks had discrimination scores similar to NH, perhaps indicating these individuals may benefit from musical training. Onset responses and acoustic change complexes can be recorded in response to instrument changes, although additional investigations need to be completed to correlate these responses with behavioral data.
characteristic scale in CI users was positively correlated with speech scores (CNC R=0.260; AZ Bio Quiet R=.510), although not as strongly as shown in previous studies.

**Conclusion**
The relative decline in performance when musical notes were manipulated was similar for CI users as for NH subjects. As such, information such as attack and decay are important for instrument identification, but are not excluded by the processor or implant transmission. Instead, the data suggest that additional spectral elements of timbre are not being encoded.

**Music perception associated with the quality of life in post-lingually deafened cochlear implant users**

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**Background**
The perception of music remains challenging for adult cochlear implant users. Nevertheless music is a pervasive art form, an environmental sound and a potent pleasurable stimulus that can be used to positively affect emotional state. As such, music therapies have shown to improve the quality of life for terminally-ill patients and older adults, an effect that could be relevant for CI users. Therefore in the present study we hypothesized that better enjoyment, quality, and perception of music in post-lingually deafened cochlear-implant users may be associated with higher health-related quality of life and better hearing ability.

**Methods**
Ninety-eight cochlear-implant users evaluated themselves on: 1) music enjoyment, quality, and perception (Dutch Musical Background Questionnaire), 2) health-related quality of life (Nijmegen Cochlear Implant Questionnaire) and 3) hearing ability (Speech, Spatial and Qualities of hearing Scale). In addition, percent-correct scores of word recognition in quiet were gathered.

**Results**
All music measures were correlated with the quality of life; only one music measure, the subjective perception of the elements of music, was correlated with the self-reported hearing ability. Furthermore, both quality of life and self-reported hearing ability were moderately correlated with the word perception in quiet. Multiple regression analyses showed the perception of elements of music and words to account for 34\% of the variance in quality of life and the perception of elements of music and words and the music listening habits for 46\% of the variance of the hearing ability.

**Conclusion**
In support of our hypothesis all music measures were correlated with the quality of life. In partial support of our hypothesis only one music measure, the subjective perception of the elements of music, and the speech perception measures were correlated with the hearing ability. An explanation for this finding may be that both the self-reported hearing-ability and the subjective perception of the elements of music are more pointed towards measuring performance and ability than the health-related quality of life, which covers more domains of daily life. Concluding, a musical therapy program or new signal processing algorithms that enable CI users to enjoy music more may improve their quality of life and partially their hearing ability.

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**Background**

Improved signal detection (i.e., enhancement) of tones occurs when a component of a complex tone is originally absent from the signal and is inserted at some later time. While enhancement has been shown to occur in normal-hearing (NH) and cochlear-implant (CI) listeners, it has yet to be shown to occur in hearing-impaired (HI) listeners. We therefore compared enhancement in NH, HI, and CI listeners under conditions of varying stimulus level and spectral spacing.

**Methods**

Ten NH, six HI, and six unilateral CI listeners participated in the study. Detection thresholds were measured with an adaptive procedure for a component in a 100-ms complex tone (test stimulus). Enhancement was defined as the difference in the thresholds between the test stimulus preceded by a 1-s adapter and the test stimulus alone. NH and HI listeners were presented harmonic complexes with a fundamental frequency (F0) of 100 or 200 Hz. The target frequency (TF) was 1 or 2 kHz. Spectrum levels were varied between 23 and 63 dB SPL. CI listeners were presented constant-amplitude pulse trains over multiple electrodes via direct stimulation. Electrode spacing, number of background electrodes, and level were varied.

**Results**

All three types of listeners showed enhancement, i.e., lower detection thresholds of the target component/electrode with the adapter present. For NH listeners, enhancement was independent of overall level and was largest for F0 = 200 Hz and TF = 1000 Hz. HI listeners also showed enhancement, but it was less than that observed for NH listeners, particularly in regions of hearing loss. All CI listeners showed enhancement; individual tendencies were prevalent, though most showed larger enhancement for larger levels.

**Conclusion**

Stimulus parameters (e.g., spectral spacing) as well as subject factors (e.g., hearing loss) affect the magnitude of enhancement. Comparison of enhancement across listener groups might elucidate the mechanisms necessary for enhancement to occur. For CI listeners, conditions that show enhancement may inform future processing strategies on how to improve speech understanding in noise.
Assessing the Effects of Reverberation on Musical Sound Quality Perception in Cochlear Implant Users
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Background
In previous studies, we designed a musical sound quality assessment method (called CI-MUSHRA) to quantify specific acoustic parameters that contribute to cochlear implant (CI)-mediated sound quality impairments. The purpose of this study was to assess the acoustic parameter, reverberation time (RT60; time for sound to decay by 60dB), using CI-MUSHRA. It is well established that acoustic reverberation plays an important role in the perceived sound quality of music by adding fullness and a sense of space; without reverberation, music sounds dry and lifeless. We hypothesized that the difficulty of CI users to perceive reverberation cues contributes to their overall musical sound quality impairments.

Methods
To determine the appropriate stimuli for the CI-MUSHRA evaluation, 6 classical musical pieces (solo and ensemble) that were originally recorded using an anechoic sound chamber were modulated with various RT60s (0, 1, 2, 3, 4, 5, or 6 seconds). 9 normal hearing (NH) listeners completed a two-alternative forced choice (2AFC) paradigm in which they were presented with two RT60 versions of a musical piece and asked to select the stimulus in the pair with the best sound quality.

For the CI-MUSHRA evaluation, five RT60 versions of a classical piece (0, 1, 2, 4, 6 second RT60 versions) were simultaneously presented in random order, along with a labeled-reference (2 second RT60; best sound quality version as rated by NH listeners). One CI participant listened to each version and provided sound quality ratings based on a 100-point scale that reflected perceived sound quality difference among versions. CI user data accrual is ongoing.

Results
For the 2AFC paradigm, NH listeners rated pieces with small amounts of reverberation (1-2 second RT60) as having better sound quality than anechoic versions (0 second RT60) and versions with greater than 2 second RT60 (Bradley-Terry Model). These results revealed that small amounts of reverberation play an important role in perceived sound quality. In comparison, for the CI-MUSHRA evaluation, the one CI user tested could not detect sound quality difference among most RT60 versions (p>0.05; two-way ANOVA), suggesting that difficulties perceiving reverberation cues contributes to implant-mediated sound quality impairments.

Conclusion
These preliminary results indicate that the impaired ability of CI users to perceive reverberation cues may contribute to musical sound quality limitations. Processing strategies that can enhance perception of acoustic reverberation have the potential to improve implant-mediated music listening.

Effect of site-specific level adjustments on speech recognition with cochlear implants
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Background
Modulation detection thresholds (MDTs) vary across stimulation sites in a cochlear implant electrode array in a manner that is ear specific. Previous studies have demonstrated that speech recognition with a cochlear implant can be improved by removing stimulation sites with poor modulation sensitivity from a subject's processor MAP. The objective of the current study was to test an alternative site-specific approach for improving speech recognition. Based on previous findings that modulation detection improves significantly with stimulus level, adjustments in level were made at stimulation sites with the poorest MDTs.

Methods
Nine postlingually-deafened ears implanted with the Nucleus cochlear implants were evaluated for MDTs, detection threshold levels (T levels) and the maximum-comfortable loudness levels (C levels) on each of the available stimulation sites. In six ears, these measures were compared to measurements taken on average 1.4 years prior to the study to test for stability over time. For each ear, the minimum stimulation level settings in the speech processor MAP were raised from true thresholds (i.e., Ts) on 5 stimulation sites with the poorest MDTs. Level increases of 5% and 10% of the dynamic range were tested. For comparison, a 5% level raise was applied to all stimulation sites. Speech reception
thresholds using the level-adjusted processor MAPs were compared to a control MAP that utilized true threshold settings without level adjustment.

Results
In most cases, the across-site patterns of Ts, Cs, and MDTs were consistent with those measured 0.9 to 1.8 years earlier. On average, the 5% level increase on the 5 electrodes with the worst MDTs resulted in an across-ear mean improvement in speech reception thresholds of 2.36 dB relative to the control MAP. For individual ears, the 5% raise almost always produced better performance relative to the control. Performance with the 10% raise produced the best performance in 3 of the 9 ears but performance was equal to or less than that with the 5% raise in 4 ears and it was worse than the control in 2 ears. The 5% level increase on all stimulation sites, a parsimonious approach, was not effective compared to the site-specific level adjustments.

Conclusion
Across-site patterns of psychophysical measurements remained stable in experienced implant users over a test-retest period averaging 1.4 years. Level adjustment was an effective alternative to site selection for improving speech recognition in cochlear implant users.

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Sound-field Auditory Steady State Response (ASSR): Data from Normal-Hearing Adults, Cochlear Implant Recipients and Cochlear-Implanted Human Cadaveric Model

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Background
Objective sound-field (SF) techniques need to be explored for evaluating difficult-to-test cochlear-implant (CI) recipients as they may provide real-world information not provided by electrical pulses. The purpose of the present study was:

Experiment 1: To determine if the SF-ASSR can be measured for adults with normal-hearing (NH) and CIs. To investigate if electrophysiologic ASSR thresholds (E-ASSR) obtained through SF correspond to the subject’s behavioral thresholds to both narrow band noise (NBNthds) and ASSR stimulus (B-ASSR).

Experiment 2: To evaluate E-ASSR data for artifact that may need to be controlled for in this and future studies.

Methods
Experiment 1: Subjects included 15 adults with NH and 6 adults with CI. ASSR measurements (IHS SmartEP) were made with a SF speaker located 1 meter from test-ear and at 90° Azimuth, with the non-test ear plugged. Recordings were made from two montages: forehead to contralateral-mastoid and forehead to nape, ground: contralateral clavicle. Thresholds were obtained for both groups in three conditions: NBNthds, B-ASSR, and E-ASSR for 0.5, 1, 2, 4.0 kHz.

Experiment 2: The right ear of a human cadaver (CIHC) was implanted with a CI 6 hours post-mortem. Normal impedances indicated that the CI was placed in an electrically conductive environment, while Neural Response Telemetry was absent for all channels. ASSR was recorded with the same protocol outlined in experiment 1.

Results
Experiment 1: Statistically significant differences between NBNthds, B-ASSR and E-ASSR were present for the NH group. A subtraction factor could be applied to predict behavioral thresholds if E-ASSR is known. There were no significant differences between the three conditions for the CI group (Figure1) suggesting that E-ASSR may be a good predictor of behavioral thresholds.

Experiment 2: In the CIHC, ASSR was recorded despite nonexistent neural activity (Figure2), suggesting that the artifact “mimicked” the response. Across frequencies, as intensity of the stimulus increased from 40 to 70 dB SPL, the amplitude of the artifact increased irrespective of the site of recording (contralateral mastoid vs. nape). Also, when the stimulus characteristics were held constant, amplitude of the artifact was greater when recorded from the contralateral mastoid compared to the nape of the neck.

Conclusion
Further data is needed to characterize and control the artifact, which will enhance the clinical applicability of SF-ASSR, especially in assessing difficult-to-test CI recipients.
Speech enhancement using sparse coding in a neural space
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Background
In normal hearing, the ability to understand speech is remarkably robust even in the presence of high levels of noise and reverberation. In contrast, hearing-impaired listeners, including those with cochlear implants, have great difficulty understanding speech and following a conversation even when the speech is considerably more intense than the noise. Similarly, automatic speech recognition algorithms are highly susceptible to corruption by interfering noise of even moderate intensity. The superiority of human listening motivates a neural approach to speech processing. Sang et al (2011) proposed just such an approach, based on the hypothesis that the response of each class of neuron in the ventral cochlear nucleus can be modeled by fractional derivatives of order k, where \(0 \leq k \leq 1\). This provides for a multi-dimensional neural space in which speech sounds can be represented. An efficient representation of speech such as this may be one factor contributing to the robustness of human speech perception.

Methods
Employing a sparse-coding strategy, we tested the potential of using a multi-dimensional neural space as a biologically-inspired basis for noise reduction and speech enhancement. Noisy speech was processed with an auditory model and the output of each frequency channel transformed into a neural space with ten k dimensions. Sparse coding was performed on the speech in the neural space and the enhanced speech signal was then reconstructed.

Results
Informal listening indicated that the level of the background noise had been reduced. The intelligibility of the reconstructed enhanced speech will be formally tested in normal hearing listeners using the BKB sentence corpus. The results of these experiments will be reported at the conference.

Conclusion
Fractional derivatives enable transformation of speech signals into the neural space, and these can be resynthesized without loss of quality. De-noising can be performed by sparse coding on fractional derivatives and may assist noise reduction for hearing prostheses and automatic speech recognition.
Radiant Energy, Pulse Shape, and Power as Driving Parameters for Infrared Neural Stimulation in the Cat Cochlea

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Background
Infrared neural stimulation (INS) is a non-contact mode of spatially selective stimulation that shows promise for future applications in cochlear implants (CIs). Delivery of radiation pulses results in heat deposition. It is crucial to optimize the stimulation parameters to prevent tissue damage.

Methods
We conducted physiological studies of the stimulus parameters (radiant energy, pulse duration, and pulse shape) in cat cochleae using the Capella, Renoir, and RINS-Alpha lasers. The physiological response of auditory neurons was quantified by measuring evoked compound action potentials (CAPs) from the round window.

Results
Pulses of equivalent radiant energy but different pulse durations were used for INS. Shorter pulses were more effective. For pulses above approximately 100 µs, the response scaled with power. Investigations of the effect of pulse shape showed that waveforms with a rapid onset to, or rapid offset from, peak power elicited the largest responses. At low values, further reductions in radiant energy decreased the CAP response.

Conclusion
The results suggest that short duration pulses of shapes with a rapid rise to, or fall from, peak power that delivered sufficient radiant energy were the most efficient form of INS and should be utilized in the development of CIs based on INS.

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Optimization of cochlear implants coding strategies using spike distance metrics and information theory

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Background
Over the past 20 years, different cochlear implant coding strategies have been proposed to improve the patients' speech recognition performance. Frequently, vocoders are used on normal hearing listeners (NH) as a model to study the impact of specific psycho-acoustical effects (e.g. dip-listening in co-modulation masking release) or new coding strategy (e.g. strategies using both amplitude and frequency modulation). A major problem of this approach is to generate stimuli carrying identical information to NH and CI listeners: place coding (that corresponds to most common case of envelope coding by the cochlear implant) is accurately simulated, however fine temporal coding cannot be simulated through vocoder technique. In addition, the vocoded stimuli are ultimately re-analysed by the healthy ear with its distortion products and no peripheral degeneracy.

Methods
We use a computational model of electrical stimulation to simulate the responses of auditory nerve fibers (ANF) to electric stimulation. The model includes the CI pre-processing stages, computation of the intra-cochlear potential and its effect on a biophysical model of ANF. First, we fit the model to the results of Miller et al. (2001) before optimizing pre-processing parameters using two different cost functions. In both cases, the cost functions are related to the distance between spike trains produced by the ANF population. We use a two parameters bin-less spike distance metric (Aronov 2003) where the first parameter describes the cost of moving spikes in time while the second that of swapping discharge across cells.

Results
We present a first cost function which measures the distance between spike trains produced by electrical stimulation and normal responses to acoustic sounds. Responses are simulated following presentation of vowel-consonant-vowel (VCV) at different signal-to-noise ratio (SNR). The second cost function solely uses responses to electrical stimulation. The optimization is done by assessing the information carried by spike trains encoding the finite set of VCV sounds. We show that the temporal and spatial resolution at which distances are computed can have a major effect on the pre-processing parameters. We discuss the results in relation with tolerance to electrode placement and peripheral nerve survival.

Conclusion
This study provides a new quantitative method for improving coding strategies based on the peripheral information supplied by CI.
An objective criterion for detecting threshold of evoked potential based on a likelihood ratio test.
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Background
Electrically evoked compound action potentials (ECAP) are routinely used in clinical audiology to set the threshold level of a patient’s cochlear implant (CI) processor. In comparison to psychophysical experiments where threshold are usually estimated using an adaptive procedure, the estimation of threshold from evoked potentials is often left to the judgment of experts. Only a few statistical tests have been proposed to set objective criteria for the detection of an evoked potential. For instance, Hughes et al. (2000) propose to compare a template to the recorded response. Threshold is set when a minimal level of correlation with the template is obtained. More recently, Undurraga et al. (2012) proposed a criterion for threshold estimation based on the deviation from F-statistics between the signal and residual noise measured. In this study, we demonstrate the use of a threshold criterion based on a likelihood ratio test.

Methods
First, numerical simulations are carried to demonstrate the validity of the likelihood ratio test in setting a threshold for signal detection. The simulation use a template signal embedded into a Gaussian white noise of different levels. Second, electrically evoked compound action potential (ECAP) recorded from implanted guinea-pigs are analysed. Animals are implanted using miniaturized electrode array and stimulated by biphasic current pulses. A forward masking paradigm is used for artifact subtraction to obtain the ECAP.

Results
We demonstrate both analytically and using simulation that the likelihood ratio test is equivalent to the F-statistic in the case of a one-sample signal. We then show that prior knowledge of the signal shape can be used to decrease the minimal SNR for detection. We investigate the effect of filtering on the recordings and show how to derive optimal filter parameters for a signal-filter pair. Finally, we validate the approach on ECAP recordings obtained from anethetised guinea-pig and discuss existing drawbacks linked to partial artifact cancellation.

Conclusion
In conclusion, using a standard approach in signal detection theory, we can provide a statistical test allowing to decide on the presence or absence of an evoked response.

Enhanced Temporal Coding in Auditory Prostheses of Onsets in the Speech Envelope can lead to Improved Sound Perception in Adverse Listening Conditions
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Background
Speech intelligibility (SI) can be very good in quiet for users of hearing aids (HAs) or cochlear implants (CIs). In adverse listening conditions their speech understanding rapidly decreases. Recent studies have confirmed that the most important parts of the speech signal for SI are the phoneme transitions and the rapid changes in temporal and spectral content. In this research a speech enhancement algorithm called enhanced envelope (EE) strategy has been developed that focuses on the enhancement of the onsets of the speech envelope. The strategy amplifies the onsets of the speech envelope in all frequency bands by deriving and introducing additional peak signals at the onsets.

Methods
The algorithm was evaluated with two groups of hearing impaired listeners: HA and CI users. The potential of the EE strategy was investigated in stationary speech shaped noise (SSN) and in an interfering talker. Keyword correct scores were collected for the Dutch LIST sentences in stationary SSN with six CI users at fixed signal-to-noise ratios (SNRs). Additionally, the speech reception threshold (SRT) was determined in an adaptive procedure in stationary SSN and in the competing talker scenario with 5 HA and 5 CI users, respectively. The presentation level of the speech was 65 dB SPL. For the HA users, the presentation level was fitted to the hearing loss using NAL-rules.

Results
All CI users showed an immediate benefit of the EE algorithm in comparison to the reference strategy that they were using in their clinical device. Overall, an SRT improvement of 2 dB was obtained in stationary SSN and around 3 dB in the
competing talker condition when the onsets of the target speaker were enhanced. Furthermore, a benefit was obtained when the onsets of the noisy mixture were enhanced. The latter does not require a priori knowledge of the speech signal for the peak signal extraction. In HA listeners, an SRT improvement of 1.5 - 2.5 dB was obtained in stationary SSN.

Conclusion
The results suggest that SI can be improved in adverse listening conditions by an enhanced temporal coding of the onsets of the speech envelope in the signal processing path of auditory prostheses.

A Case Study of Optimized Cochlear Implant Stimulation in a Congenitally-Deaf Individual
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Background
In post-lingually deafened individuals, cochlear implants (CIs) are effectively intended to achieve the ideal condition of replicating excitation patterns in the auditory nerve produced under normal acoustic hearing conditions. Under this ideal, perception of external acoustic stimuli would be unchanged relative to performance prior to onset of deafness. However, few efforts have been directed at identifying the upper-bound of performance that is achievable with current CI hardware and signal processing. Rather, current stimulation patterns have often been developed in a phenomenological manner. We have developed a framework by which CI stimulation may be optimized for an individual, and documented substantial performance benefit in a vowel identification task.

Methods
In our framework (Figure 1), an acoustic source is passed through (top) a computational model of normal acoustic hearing (Heinz et al., ARLO, 2001) and (bottom) a cochlear implant electrode selection algorithm followed by a computational model of electric hearing (Bruce et al., IEEE TBME, 1999), yielding two neural activation patterns (NAPs). These NAPs are compared using an objective function to obtain a distance measure that is used to select electrodes to best match the NAP produced by the model of electrical hearing with the NAP produced by the model of normal acoustic hearing. This optimizes the electrical stimulation sequence against the nerve activity expected to be input to the central auditory pathway in the case of a healthy cochlea.

Optimized stimulation patterns were generated for a 23 y.o. congenitally-deaf user (14 years; Nucleus-24). Patterns were generated for nine /hVd/ vowel tokens, each presented in quiet (infinite SNR) or speech spectrum-shaped noise (20 and 10 dB SNR). The Nucleus MATLAB Toolbox drove the user’s implant while performing a 9AFC vowel identification task under his preferred (real-time) and optimized (pre-generated) strategies, using an A-B-A-B (by day) test paradigm.

Results
Substantial benefit was evidenced for optimized stimuli (Figure 2) under conditions of speech in quiet and noise.

Conclusion
Appreciable improvement in a subject for whom this optimization is non-ideal (a congenitally-deaf user is not expected to better recognize "normal acoustic hearing" inputs) suggests that CI stimulation patterns can be improved to increase speech perception performance, even in modest background noise. Use of this framework can provide a lower-bound on the ceiling for CI speech perception performance, against which future improvements in hardware and stimulation strategy may be measured.
Asymmetric Signal Degradation Causes Difficulty in Understanding Speech in Spatial Attention Tasks
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Background
Normal-hearing (NH) listeners can nearly perfectly recall words in an attended ear while ignoring the other ear when dichotic speech is presented over headphones. However, recent experiments have shown that this does not necessarily occur in people with asymmetrical hearing. People with asymmetrical hearing loss, bilateral cochlear implants, or single-sided deafness very likely have a relatively “good” ear (i.e., high-quality speech signals) and “bad” ear (i.e., low-quality speech signals). We simulated different levels of signal degradation by independently varying the signal quality in each ear, as would occur in users of one or two cochlear implants, in order to understand the effect of having a good and bad ear in spatial attention tasks.

Methods
Ten NH listeners participated in the study. Listeners reported keywords from five-word nonsense sentences in selective and divided attention tasks. Over headphones, a target talker was presented in one ear and an interferer was presented in the other, both at 65 dB SPL-A. In experiment 1, signal degradation was imposed by varying the number of channels in a noise vocoder. The signals were either unprocessed speech or had 4, 8, or 16 channels. In experiment 2, signal degradation was imposed by tonotopically shifting carrier frequencies in an eight-channel sine vocoder. Training was provided prior to testing for the shifted sine-vocoded sentences.

Results
For experiment 1, in conditions with symmetrical signal degradation, in the selective attention task, listener performance was near ceiling for all conditions; hence, there was no effect of the number of channels. In the divided attention task, performance was near 50% correct on average and gradually increased as the number of channels increased. In conditions with asymmetrical signal degradation, in both the selective and divided tasks, listener performance was systematically lowest when there was poorer resolution in the attended ear compared to the ignored ear, and was highest when there was better resolution in the attended ear compared to the ignored ear. It was hypothesized that a similar trend would be seen in experiment 2 when tonotopic shift was imposed.

Conclusion
These data demonstrate that it is more difficult to attend to a poor-quality speech signal when trying to ignore a high-quality speech signal. This may adversely affect the advantage of spatial listening in certain spatial configurations for people with asymmetrical hearing, i.e., those with good and bad ears.
Vertical Perception of Simple Stimuli in Normal-Hearing and Cochlear-Implant Listeners
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Background
In normal-hearing (NH) listeners, elevation perception depends upon frequency-dependent peaks and notches imposed by filtering from the pinna. However, limited spectral resolution and current spread in cochlear-implants (CIs) make it difficult to encode spectral cues. Previous research has shown that elevation can be perceived from single tones. Sounds presented over headphones or with CIs lack filtering from the pinna and are perceived “within” the head. We examined the perceived intracranial elevation of tones in NH listeners and single-electrode stimulation in CI listeners. We hypothesized that post-lingually deafened CI users would show systematic changes in elevation with electrode number, while pre-lingually deafened CI users would not. The ultimate goal was to determine the importance of typical elevation perception development from acoustic stimulation.

Methods
NH listeners were presented monaural and diotic tones varying from 0.125 to 16 kHz (19 frequencies) using ear-insert headphones. Stimuli were calibrated to a comfortable and equal loudness (reference of 50 dB SPL-A at 1000 Hz). Unilateral and bilateral stimulation was presented to two bilateral CI listeners; unilateral stimulation was presented to three unilateral CI listeners. Stimuli were biphasic, monopolar pulse trains at one of 13 electrodes or electrode pairs, all presented at comfortable levels. The rate of pulse trains was 300 or 1000 pulses per second (pps); conditions with rate roving were also tested (50% randomization) to confound assigning elevation simply to pitch. On each trial, listeners indicated where they heard the auditory image by clicking on a face (horizontal and vertical position).

Results
Preliminary NH data showed that listeners perceive a change in elevation as frequency increases. Specifically, sounds are perceived higher as the frequency increases between 0.25-12 kHz and does not change at higher frequencies. Most post-lingually deafened CI users showed that the basal electrodes were often perceived as higher in elevation and responses were unaffected by rate roving. No prelingually-deafened CI users showed a change in elevation that was unaffected by rate roving, implying that these listeners were simply assigning elevation to pitch.

Conclusion
These data highlight the association between place of stimulation and elevation perception in NH and CI listeners. Post-lingually deafened individuals may be making use of their spatial maps developed prior to hearing loss and may help us understand how the auditory system develops the perception of elevation. These data also demonstrate the potential to present elevation cues to CIs.

ACOUSTIC MODELS OF COCHLEAR IMPLANTS: EVALUATION IN SINGLE SIDED DEAFNESS PATIENTS
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Background
Acoustic models of cochlear implants have been used in numerous published studies but the validity of these models has not been well established. A recent development presents a unique opportunity to validate and enhance these acoustic models: the implantation of “single-sided deafness” listeners, who have normal or near normal hearing in the ear contralateral to the implant. These listeners can compare what they hear with the implant to what they hear with an acoustic model in the normal hearing ear.

Methods
Three cochlear implant users with single sided deafness were presented the same sentence twice. First, the unprocessed sentence was sent through a direct connection to their cochlear implant speech processor (which had the microphone input deactivated). Then the sentence was processed through a cochlear implant acoustic model (a noise vocoder) and presented to the normal hearing ear through loudspeakers. The analysis filters used in the acoustic model had the same cutoff frequencies as those in the listener’s speech processor, but the noise bands of the vocoder were adjustable. Listeners used a graphical interface to determine a “self-selected model” by modifying the low- and high-frequency edge of the set of noise bands until the noise vocoder sounded as similar as possible to the percept provided by the implant. The process was then repeated for a sinewave vocoder. Lastly, listeners answered questionnaires about
the similarity between the sentences as heard through the cochlear implant and through four different types of vocoders: noise vs. sinewave, and standard vs. self selected. The standard models (which are the most commonly used in the literature) used noise bands or sinewaves whose center frequencies were identical to those of the analysis filters.

Results
Self-selected models tended to have higher frequency ranges than the standard models. On average, these self-selected models were deemed to produce sounds that were "very similar" to those heard through the implant (6.7 in a 9 point scale). Ratings for standard models were poorer on average, with a 4.7 score (where 5 corresponds to a "somewhat similar" rating. Some standard models received ratings of 3 (not very similar) and even 1 (not at all similar-completely different). Noise vocoders were better for two of the listeners and sinewave vocoders were better for the third one.

Conclusion
Results suggest that some of the acoustic models commonly used in the literature are inaccurate representations of the percepts actually heard by some cochlear implant users.

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Restriction of Wnt signaling in the dorsal otocyst determines semicircular canal formation in the mouse embryo
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Background
Inner ear develops from single epithelial sheet, otic placode. Whereas canonical Wnt signaling plays an important role in induction of the otic placode and its axial determination, its contribution to otic morphogenesis at later stages has remained unclear. We therefore have studied the role of Wnt signaling in that stage.

Methods
To investigate the expression pattern of Wnt signaling in the later stage of otic development (E10.5~E12.2), we used a Wnt reporter strain, BAT-gal. We next used a combination of whole-embryo culture and chemical compounds in order to evaluate the function of Wnt signaling.

Results
The activity of Wnt signaling was continuously observed, but gradually restricted in the dorsolateral otocyst. The genes for several Wnt ligands were expressed in the dorsal otocyst according to specific patterns, whereas those for secreted inhibitors of Wnt ligands were expressed exclusively in the ventral otocyst. With the use of whole-embryo culture in combination with potent modulators of canonical Wnt signaling, we found that forced persistence of such signaling resulted in impaired formation both of the lateral outpocketing and of the fusion plates of the dorsal outpocketing. Canonical Wnt signaling was found to suppress the expression of otic determinants, Netrin1 and Otx1.

Conclusion
Our stage-specific functional analysis suggests that strict regulation of canonical Wnt signaling in the dorsal otocyst is required for the process of semicircular canal formation.
Effects of Small RNA Depletion on Development of Mammalian Inner Ear and Sensory Hair Cell Differentiation and Maintenance

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Background
Small RNAs, namely endogenous siRNAs and miRNAs, are crucial regulators of transcriptional and post-transcriptional regulation in eukaryotic cells. Prior studies of Dicer¹ conditional knockout (CKO) effecting small RNA depletion and Dgcr8 CKO effecting specifically canonical miRNA depletion have demonstrated differences regarding cell proliferation and differentiation. While Dicer¹ CKO in stem cells has effects on both proliferation and differentiation, Dgcr8 CKO primarily effects differentiation. These results largely reflect specific roles for siRNAs in chromosomal maintenance and miRNAs in regulatory pathways effecting cell fate and function.

Methods
To compare loss of small RNAs vs. loss of canonical miRNAs in mouse inner ear, we generated Dicer¹ CKO and Dgcr8 CKO in the otic placode using Pax2-Cre and in hair cells using Atoh1-Cre. In addition to gross morphological observation, in situ hybridization was used to confirm miRNA depletion, and immunohistochemical detection of Phalloidin and MyoVIIa was used to examine development and maintenance of hair cells in each of these models.

Results
Our study of Dicer¹ CKO in mouse inner ear has shown considerable defects in gross development and maintenance of sensory hair cells depending on the timing and tissue specificity of conditional deletion. We hypothesized that Dgcr8 CKO would result in less severe outcomes. To the contrary, preliminary analysis of Dgcr8 CKO demonstrates that depletion of only canonical miRNAs results in greater variability of developmental outcomes in the Pax2-Cre model and surprisingly more detrimental effects on sensory hair cell differentiation and maintenance in the Atoh1-Cre model.

Conclusion
Results are interpreted to reflect the role of miRNAs in canalization (i.e. reproducibility of developmental outcomes) and the importance of the balance of small RNA functions in cell differentiation and maintenance.

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Background

Mutations in PCDH15 gene can cause Usher syndrome type 1F (USH1F) in humans, characterized by congenital deafness, vestibular dysfunction and progressive retinal degeneration. Animal models harboring mutations in the PCDH15 gene (i.e. Ames waltzer and orbiter) show different degrees of splaying stereocilia reduce FM1-43 uptake and abnormal transducer currents, suggesting the involvement of PCDH15 in hair bundle development and function. A role in synaptic maturation has also been suggested for specific PCDH15 isoforms. Zebrafish have duplicate pcdh15 genes, pcdh15a and pcdh15b. Previous observations from mRNA in situ hybridization studies suggested the existence of specific and unique roles for Pcdh15a and Pcdh15b in the ear and eye, respectively.

Methods

We developed antibodies directed to the more divergent intracellular tails of Pcdh15a and Pcdh15b. Western blot analysis and immunohistochemistry experiments demonstrated the expression of both Pcdh15 proteins in ear hair cells from 1 day post-fertilization (dpf) to 5dpf.

Results

In hair cells Pcdh15a is restricted to the stereocilia hair cell bundle while Pcdh15b is also expressed at the hair cell bundle and below the cuticular plate. Neuromast hair cells show high expression of both Pcdh15 proteins in the kinocilia, the hair cell bundle and in the cuticular plate. Modest expression is also observed at the basolateral membrane and hair cell body. In the zebrafish retina (4dpf) we observed expression of Pcdh15b only at the connecting cilia in photoreceptor cells. No expression was observed for Pcdh15a in zebrafish eye under our experimental conditions. These last observations are in agreement with previous published work showing only expression of Pcdh15b in fish retina.

Conclusion

Taken together, the results presented here show for the first time the spatiotemporal pattern of expression of both Pcdh15 proteins in zebrafish and also may explain the mild splayed stereocilia phenotype and the normal planar polarity observed in orbiter mutants where only Pcdh15a is mutated.

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Ethanol Affects the Development of Sensory Hair Cells in Larval Zebrafish (*Danio rerio*)

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**Background**

Some children who are born to mothers who have consumed alcohol during pregnancy present a number of morphological, cognitive, and sensory abnormalities, including hearing deficits, collectively known as fetal alcohol syndrome (FAS). The goal of this study was to determine if the zebrafish lateral line could be used to study sensory hair cell abnormalities caused by exposure to ethanol during embryogenesis.

**Methods**

Lateral line hair cells are present at 2 days post-fertilization (dpf) and by 5 dpf sensory hair cells are functional. We treated a transgenic line of zebrafish that expresses membrane-targeted green fluorescent protein under control of the Brn3c promoter/enhancer with embryo medium supplemented with varying concentrations of ethanol (0.75%-1.75% by volume) from 2 dpf through 5 dpf. Age-matched, non-treated controls were also raised in parallel. The vital dye FM 1-43FX was used to detect the presence of functional mechanotransduction channels. Larval zebrafish were fixed, mounted, and imaged using a confocal microscope.

**Results**

As the concentration of ethanol increased, we observed some classical abnormalities associated with FAS; enlarged heart, small eye, and craniofacial defects. Moreover, the number of sensory hair cells decreased as the concentration of ethanol increased in a dose-dependent manner. Also, the percentage of FM 1-43-labeled hair cells decreased as the concentration of ethanol increased.

**Conclusion**

Ethanol treatment of zebrafish embryos during the time when hair cells are added to the lateral line not only results in fewer hair cells, but those hair cells that are formed do not function properly. Our results mirror data obtained by investigators who have analyzed the effect of ethanol on auditory function in developing mammals. Zebrafish may prove to be a useful model system to study the mechanisms underlying the development of FAS.

Roles of Primary Cilia in Mammalian Inner Ear Development

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**Background**

Primary cilia are microtubule-based structures that extend from the surface of all mammalian cells. These non-motile cilia serve as a cellular signaling center for many pathways including Sonic hedgehog (Shh), Wnt, FGF, and PDGF. Defects in the formation or function of primary cilia are shown to be associated with a range of genetic diseases collectively known as ciliopathy. Hearing loss is one of the well-characterized phenotypes found in human ciliopathy, yet the underlying pathological mechanisms remain unclear.

**Methods**

In order to understand the role of primary cilia in inner ear development, we analyzed the inner ears of a ciliopathy mouse model, in which ciliary morphology is defective due to a missense mutation in *Broad minded* (*Bromi*) encoding an uncharacterized TBC (Tre-2, Bub2, and Cdc16) domain-containing ciliary protein.

**Results**

We observed that inner ears of *Bromi* mutants exhibited diverse inner ear defects, some of which are attributed to defective Shh signaling. For example, the cochlear duct of *Bromi* mutants was severely shortened and lacked the expression of *Msx1*, an apical cochlear marker, suggesting a failure of apical cochlear specification that requires high levels of Shh signaling. Multiple extra rows of hair cells (HCs) were observed in the organ of Corti in the apical cochlea, suggestive of a failure of convergent extension of the prosensory domain. However, HC orientation appeared unaffected in *Bromi* mutants based on normal localization of the kinocilia at the abneural side of HCs. Interestingly, ectopic sensory patches containing vestibular-like HCs developed in the Kolliker's organ in *Bromi* mutants, similar to phenotypes reported in *Gli3*Δ699/Δ699 mutants, in which Shh signaling is compromised. Based on *Atoh1* expression and hair bundle morphologies, HCs were more mature in *Bromi* mutants, consistent with the in vitro results of Shh inhibiting HC formation. Additionally, formation of primary cilia was severely affected in most of the supporting cells of *Bromi* mutants but not in the
HCs, consistent with a previous report that *Bromi* mutants display neural tube or limb defects associated with abnormal cellular responses to high levels of Shh signaling.

**Conclusion**

Our results suggest that the phenotypes observed in *Bromi* mutants are due to compromised Shh signaling and that Shh signaling in the mammalian cochlea is mediated through the primary cilia.

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**Spontaneous activity in auditory inner hair cells drives ribbon synapse maturation during a critical period**

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**Background**

The maturation of neural circuits is thought to rely on spontaneous electrical activity that occurs during immature stages of development (Katz & Shatz, 1996 Science 274: 1133-8; Blankenship & Feller, 2010 Nat. Rev. Neurosci. 11: 18-29). In the developing mammalian auditory system, the sensory inner hair cells (IHCs) fire calcium action potentials (Tritsch et al. 2007 Nature 450: 50-5; Johnson et al. 2011 Nat. Neurosci. 14: 711-7). IHC action potentials are generated by the interplay between a depolarizing Ca²⁺ current and a repolarizing delayed rectifier K⁺ current (Marcotti et al. 2003 J. Physiol. 548:383-400). However, the ability to sustain repetitive firing activity depends largely upon the activation of the transiently expressed small conductance Ca²⁺-activated K⁺ current SK2 (Johnson et al. 2007 J. Physiol. 583: 631-46).

**Methods**

Using a transgenic mouse in which the repolarizing potassium current, SK2, is normally over-expressed, but can be down-regulated in vivo using doxicycline (Hammond et al 2006 J. Neurosci. 26: 1844-53), we investigated whether normal spiking activity is required for IHC development. Electrophysiological experiments were carried out using physiological conditions (body temperature and 1.3 mM extracellular Ca²⁺). Whole-cell patch clamp was used to measure current and voltage responses in immature and adult IHCs and their exocytotic properties, which were assessed by measuring changes in cell membrane capacitance (ΔCm).

**Results**

We found that the larger SK2 current in immature IHCs of SK2 over-expresser (SK2-OE) mice made the action potential hyperpolarization phase more robust, which consequently reduced spike width and increased their frequency compared to controls. The consequence of abnormal immature spontaneous activity was that SK2-OE IHCs failed to develop the linear exocytotic Ca²⁺ dependence that is a characteristic of adult cells (Johnson et al. 2010 Nat. Neurosci. 13: 45-52), indicating a direct role for spontaneous Ca²⁺ action potentials in regulating IHC maturation. Using the ability to down-regulate the over-expression of SK2 channels in OE mice with doxicycline, we were able to define the “critical period” over which this regulation occurred.

**Conclusion**

Our results provide the first direct evidence that maturation of mammalian primary auditory receptors depends upon a specific temporal pattern of spontaneous calcium signals that occurs during a critical period prior to the onset of sound-driven neuronal activity.

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Conditional deletion of *Pten* leads to defects in the spiral ganglion by modulating Akt/GSK3β signaling in the inner ear

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Background

All cellular phenomena and developmental events, including inner ear development, are modulated through harmonized signaling networks. Phosphatase and tensin homolog deleted on chromosome 10 (PTEN), a tumor suppressor, is a major signaling component involved in crosstalk with key regulators of development, i.e., Wnt, Notch, and bone morphogenetic proteins. Although *Pten* function has been studied in various systems, its role in inner ear development is poorly understood.

Methods

Here, we used inner ear-specific *Pten* conditional knockout mice and examined the characteristics of the inner ear.

Results

In a detailed analysis of the phenotype, reduced cochlear turning and widened epithelia were observed. Phalloidin staining of sensory epithelium revealed that hair cell patterns were disturbed, i.e., additional rows of hair cells were discovered. The neural abnormality revealed a reduction in and disorganization of nerve fibers, including apoptosis at the neural precursor stage. *Pten* deficiency induced heightened phosphorylation of Akt at Ser473. The elevation of inhibitory glycogen synthase kinase 3β Ser9-phosphorylation (pGSK3β) was sustained until the neuronal differentiation stage at embryonic day 14.5, instead of pGSK3β downregulation.

Conclusion

This is the first report of the influence of *Pten/Akt/GSK3β* signaling on the development of spiral ganglia. These results suggest that *Pten* is required for the maintenance of neuroblast number, neural precursors, and differentiation in the inner ear.
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Background
Spiral ganglion cells (SGCs) play a very important role in auditory function to transmit sound signals from the cochlear hair cells to the brain. The molecular mechanism of development of SGCs has not been elucidated enough. As with other neurons, SGCs pass through mitotic proliferation, ceasing proliferation, and differentiation at a development. We focused on the Tis21/BTG2/PC3 (Tis21) gene. A Tis21 gene belongs to the member of BTG/Tob family with antiproliferative properties. Tis21 has function to induce the neuronal differentiation at cerebellum in mice (Canzoniere, Neurosci., 2004). We have already revealed that Tis21 expresses in the SGCs in embryonic stage (Hayashida, Neuroreport, 2010). We set out to perform further developmental analysis utilizing Tis21 knock-in mice.

Methods
Normal mice (Tis21(+/-) mice), heterozygous Tis21-GFP knock-in mice (Tis21(+/-GFP) mice) and homozygous Tis21-GFP knock-in mice (Tis21(-GFP/-GFP) mice) were used for this experiment. We performed histological analysis of GFP expression in Tis21(+/-GFP) mice and Tis21(-GFP/-GFP) mice, and we counted the number of SGCs of Tis21(-GFP/-GFP) mice and Tis21(+/-+) mice.

Results
The GFP expression was detected in the cytoplasm of SGCs at Embryonic day15.5 (E15.5), E18.5 and Postnatal day4 (P4) in Tis21(+/-GFP) mice and Tis21(-GFP/-GFP) mice. The number of SGCs in Tis21(-GFP/-GFP) mice was significantly decreased when compared with that of Tis21(+/-+) mice at E15.5, E18.5 and P4.

Conclusion
Tis21 (-GFP/-GFP) mice showed significant decrease of number of the SGCs. This finding suggestd that Tis21 is involved in development of SGCs. Neurogenin1(Ngn1) deficient mice show complete absence of SGCs (Ma, Otolaryngol., 2000). NeuroD deficient mice show mostly complete abscent of SGCs, primarily in the apex. It is considered that Ngn1 acts upstream of NeuroD during development of SGNs (Liu, Genes Dev., 2000). In additon, Tis21 acts upstream of Math1 during cerebellum developmet (Canzoniere, J Neurosci., 2004). Considering these facts, Tis21 probably acts downstream of Ngn1 during development of SGCs.

Conditional deletion of neurotrophins indicate their essential role in postnatal fiber reorganization

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Background
Two neurotrophins (Bdnf, Ntf3) support developing spiral ganglion neurons (SGNs), which degenerate in their absence. Other aspects of each neurotrophin in patterning the embryonic innervation of the ear have been partially clarified: Ntf3 is essential for the normal innervation of the organ of Corti (OC) in the basal turn of the cochlea while Bdnf plays only a supporting role for SGNs, primarily in the apex. Suggestions of the role of each neurotrophins in sorting of afferents to the two types of OC hair cells (Ntf3 for type I fibers to inner hair cells [IHCs], Bdnf for type II fibers to outer hair cells [OHCs]) were refuted by subsequent work. Early lethality has blocked analysis of the role of neurotrophins in postnatal sorting of type afferents to the two types of hair cells of the OC, leaving room for interpretations mostly based on non-experimental data. For example, some in vivo and in vitro data suggest neurotrophins may be critical for synapse formation while others suggest only a limited influence of neurotrophins on postnatal synaptogenesis and viability of neurons.

Methods
To go beyond these suggestions, we have generated viable conditional single and double mutants of both neurotrophins using hair cell and ear specific cre lines (Atoh1-cre and Pax2-cre) in which we analyze the pattern of afferent and efferent innervation and formation of synaptic contacts.

Results
We show that loss of both neurotrophins in hair cells using Atoh1-cre has only a limited effect on afferent viability but completely eliminates afferent growth to outer hair cells without influencing inner hair cells. In contrast, complete loss of either neurotrophin using Pax2-cre results in topographical disorganization of both afferents and efferents with limited innervations and synapse formation in the base in Pax2-cre; Ntf3/f mice. Finally, conditional elimination of all
neurotrophins only in the ear results in delayed loss of all afferents in postnatal mice, followed be further delayed loss of
efferents and an even further delayed loss of hair cells.

Conclusion
These data clarify for the first time in two distinct conditional deletion mouse models the role of neurotrophins in postnatal
afferent sorting to various hair cells and the role of afferents for efferent innervation to inner and outer hair cells. We show
that differential longitudinal, radial, and temporal distribution of neurotrophins define the postnatal afferent and efferent
fiber sorting to the two hair cell types of the OC.

Developmental alterations of Hair Cell Types help in Understanding the pattern of Innervation.
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Background
The organ of Corti (OC) consists of a regular mosaic of two different types of hair cells (HCs; inner hair cells, IHCs and
outer hair cells, OHCs) which are innervated in a stereotyped fashion by two types of afferents (type I to IHCs; type II to
OHCs) and efferents. In sensorineural hearing loss, HCs are lost followed by progressive degeneration of neurons.
Regaining complete hearing will require reconstituting the OC HCs with the precise sorting of their innervation. To achieve
this goal, it is necessary to understand the molecular regulation of OC patterning. Beyond generalized HC differentiation,
formation of specific types of HCs and their innervation must be understood at the molecular level.

Methods
We used different mouse models to assess the temporal and intensity variations of TF Atoh1 modulation to differentiate
specific types of HCs: Atoh1 conditional null using Pax2-cre and Atoh1-cre, Neurog1 knockin (Atoh1KINeurog1/KINeurog1), combination of Atoh1 floxed and Neurog1 KI (Atoh1cre; Atoh1f/KINeurog1), Bronx Waltzer
(bv/bv) and Pax2-cre; Neurod1f/f.

Results
The transient expression of Atoh1 in Atoh1-cre; Atoh1f/f mice result near complete loss of all IHCs. Similarly, bv/bv mutant
mice also lose IHC, with reduced Atoh1 expression at birth. In contrast, loss of Neurod1 results in OHCs specification as
ectopic IHCs due to premature and aberrant expression of Atoh1.

Molecular modifications of HCs affect innervation. In the absence of differentiated HCs in Pax2-cre; Atoh1f/f, the
radial fibers fail to reach the OC. Atoh1 KINeurog1/KINeurog1 mice can attract more fibers to the partially differentiated
OC, most likely by promoting Ntf3 expression by Neurog1. In the mutants with specific loss of IHCs display projection of
type I fibers to the remaining OHCs. In contrast, mutants with ectopic conversion of IHCs in the region of OHCs attract
type I fibers regardless of their ectopic topology.

Conclusion
We conclude that afferents can sort out IHC and OHC specific patterns if choices are possible in mutants with altered
distribution of HC types. In the absence of IHC all fibers will innervate OHCs if they cannot divert to nearby IHCs due to
increased distances to the nearest IHC. Understanding the dosage and cross-regulation of these above TFs, in regulating
specific types of HCs development and their specific fiber sorting mechanism will provide useful information for the future
attempts to restore hearing.
Countergradients and Modular Expression Patterns of Eph-Ephrin Signaling Proteins in the Developing Auditory Brainstem

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Background

A family of receptor tyrosine kinases, the Eph-ephrins, is involved in guiding topographic mapping and projection pattern formation in a variety of systems. We are interested in understanding their role in establishing the orderly arrangement of converging inputs to the inferior colliculus (IC) prior to hearing onset. Previously, we described the development of several layered projections to the central nucleus of the IC (CNIC), as well as a modular input to the lateral cortex of the IC (LCIC) arising from the lateral superior olive (LSO). Immunohistochemical studies revealed expression patterns for EphA4, ephrin-B2 and ephrin-B3 during this period of projection shaping that suggest their involvement in the mapping of these early connections. Recently, we showed in ephrin-B2 mutant mice that LSO inputs to the CNIC (bilaterally layered) and LCIC (ipsilaterally modular) lack a clear topography, despite maintaining their characteristic axonal patterns.

Methods

These results led to the present studies that build upon preliminary expression data from immunohistochemistry and X-Gal staining experiments utilizing lacZ mutants. Here, quantitative methods were used to confirm discernible CNIC gradients and LCIC expression modules, as well as to explore the possibility of similar gradients in the LSO.

Results

Sampling methods reveal gradients of EphA4 and ephrin-B2 across the tonotopic axis of the CNIC, with protein most concentrated in high-frequency regions. In addition, expression data reveals prominent modules for each throughout the LCIC. In contrast, the CNIC is devoid of ephrin-B3, although present in the LCIC and exhibiting a similar modular expression. Ephrin-B2 in the LSO is expressed in a complementary gradient, with protein most concentrated in low frequency regions. EphA4 is present in the LSO, while ephrin-B3 is absent.

Conclusion

The described countergradient between ephrin-B2 (LSO) and EphA4 (CNIC) that have known binding affinities for one another may provide insights for the establishment of topographic connections between these auditory brainstem nuclei. Further, it remains to be seen whether modular LCIC expression patterns for EphA4, ephrin-B2, and ephrin-B3 are superimposed, partially overlapping, or mutually exclusive of one another. Determining this, as well as which specific modules LSO targets, is paramount for fully understanding the role of Eph-ephrins in guiding multiple converging inputs to the IC.

The calyx of Held as a model system to study neural development: competition and pruning to mono-innervation

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Background

Several neural systems, including the neuromuscular junction and climbing fiber innervation of Purkinje cells, are considered models to study neural development because they exhibit hallmark features of initial competition and pruning, they establish a recognizable endpoint of mono-innervation of their targets and their presynaptic terminals are large and easily monitored. Several studies, using physiological counting methods and serial-section EM of relatively small numbers of cells, show evidence for competition among developing calyces of Held (CH), but its prevalence has not been properly assessed. Also, although it is well-established that the CH grows rapidly over several days during the first postnatal week, this process has not been quantified.

Methods

In order to objectively assay the dynamics of CH formation and mono-innervation of medial nucleus of the trapezoid body (MNTB) neurons, we used a primarily anatomical approach because physiological counting techniques provide an underestimate of convergent synaptic inputs due to sectioning of axons in brain slice preparations and because independent inputs may share a stimulation threshold. Serial block-face scanning electron microscopy provides high
resolution to identify neuronal contacts and the means to accurately register cellular ultrastructure through large tissue volumes.

Results
Here we make the first application of these technologies to the developing brain and study changes in neural connections across the age range critical for calyx growth (P2-P6). We establish that competition, defined as multiple large growing calyces forming synaptic contact with MNTB cells, is a common trait during this developmental period. By segmenting pre- and postsynaptic cellular structures and quantifying contact areas between nerve terminals and MNTB cells, we describe P3 as a uniquely active day in calyx growth during which cell apposition areas are added at a rate exceeding 200 μm²/day.

Conclusion
These data define metrics for growth dynamics in this system, illustrate shared features with other neural systems considered as models of neural development and exemplify utility of the CH:MNTB system for studies of synaptogenesis and pruning with relevance to other neural systems throughout the brain.

Reversal of Early Mild Hearing Loss Reveals a Discrete Critical Period for Maturation of Cortical Excitability
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Background
Sensory systems are highly susceptible to changes in peripheral activity during developmental critical periods (CP). To reveal the onset and termination of critical periods, sensory input is typically diminished or restored as a function of age. Our unpublished results in gerbils have suggested that there are at least two CPs during which deprivation influences the maturation of auditory cortex cellular properties: ear plugging immediately after hearing onset, from postnatal (P) 11-13 delays intrinsic membrane property development, while ear plugging from P14-22 disrupts voltage-gated properties. Therefore, we asked whether hearing loss induced changes to active and passive membrane properties would recover when ear plugs were removed during or after the closure of the second CP.

Methods
At P11, gerbils (Meriones unguiculatus) were implanted bilaterally with malleable plugs that induced a behavioral threshold shift of ~20 dB at 4 kHz. The plugs were removed on either P17 or P23, followed by ~6 days of normal hearing. Whole-cell current clamp recordings were then obtained from L2/3 pyramidal neurons in thalamocortical slices at P21-35 and compared to age-matched controls.

Results
When ear plugs were implanted at P11 and removed on P17 or P23, certain membrane properties eventually reached normal age-specific values (e.g., Vrest: Age-matched control = -32.5 mV ± 1.0 vs P23 plug removal = -32.4 mV ± 1.4; t = -3.6, p>0.1). Therefore, the CP for development of these properties remained open through at least P23. In contrast, specific membrane properties did not recover to normal values when earplugs were removed at P17 or P23 (e.g., AHP: Age-matched control = -14.1 mV ± 0.7 vs P17 plug removal = -12.1 mV ± 0.7; t = 0.21, p<0.05). Therefore, the CP for normal development of these membrane properties closed before P17.

Conclusion
These findings strongly suggest that even mild hearing loss can delay the functional development in cortex, and the maturation of specific membrane properties are disproportionally affected by deprivation during at least two CPs that open and close abruptly after the onset of hearing. This suggests that the restoration of normal hearing does not necessarily restore normal neuronal processing (supported by NIH DC011284 DHS and VCK).
Functional and cellular abnormalities in the developing auditory cortex after neonatal cerebral hypoxia-ischemia.

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Background
Subplate neurons (SPNs) are required for the formation and functional refinement of thalamocortical connections (Kanold & Luhmann 2010, Tolner et al. 2012). SPNs are selectively vulnerable to neonatal hypoxic-ischemic (HI) insults (McQuillen et al. 2003). Such insults in humans can give rise to auditory impairments, as well as a multitude of other developmental language disorders. Thus we investigated the consequences of HI on auditory cortex development in rodents.

Methods
HI was performed on rat pups on postnatal day (P) 1-2 and extracellular single-unit recordings were performed in anesthetized animals between P20-30 to assess the effects of HI on receptive field organization of auditory cortex neurons. Whole-cell patch clamp recordings were also performed in layer 4 of auditory cortex cells after HI on rat pups between P18-22, in order to get a more in-depth understanding of our results on a cell-specific level.

Results
We find that compared with normal controls, auditory cortex neurons in HI treated pups showed increased firing rates and abnormal tuning characteristics. Receptive fields were disrupted and thresholds were increased. Abnormal EEGs, reminiscent of seizure activity was also observed in 30 percent of the animals. After whole-cell recording we find that intrinsic properties of layer 4 cells and synaptic transmission are altered.

Conclusion
Our observations imply that SPNs are necessary for normal development of the circuit and tuning properties in the auditory cortex. Thus, injury to these neurons results in widespread abnormalities of sensory processing.
Silent synapses on subplate neurons in developing auditory cortex
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Background
Functionally silent excitatory synapses exist in several systems during development. Little is known about the presynaptic spatial origin of silent synapses. Cortical cells receive input from multiple layers and connections between these layers develop at different ages. Since cells receive inputs from multiple presynaptic cells we hypothesized that functional and silent synapses might originate in different layers.

We investigated this hypothesis in the developing circuits in auditory cortex. Subplate neurons form one of the earliest maturing circuits in the cerebral cortex and crucial to cortical development. We recently found that subplate neurons receive inputs from the developing cortical plate in the 2nd postnatal week.

Methods
We here use laser-scanning photostimulation in acute slices of auditory cortex during the 1st two postnatal weeks in mouse to study the spatial origin of silent synapses to subplate neurons.

Results
We find that silent synapses from the cortical plate can be present in subplate neurons before the emergence of functional synapses from the cortical plate. They are mostly located in the periphery of cells’ response fields and are discerned into two broad classes based on their distinct spatial patterns.

Conclusion
Since subplate neurons can be activated at these young ages by thalamic inputs, cortical neurons at young ages have the ability to signal to subplate neurons depending on the activation state of subplate neurons.

These results suggest that subplate neurons are an integral part of the intracortical circuitry at the youngest ages.
Age-related primary neural degeneration is aggravated in the absence of olivocochlear efferents
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Background
Olivocochlear (OC) efferents can reduce threshold shift from noise exposures in excess of 100 dB SPL, but the significance of this effect has been questioned, since natural environments do not typically contain such high sound levels (Kirk and Smith, 2003). We recently showed that threshold recovery after moderate-level exposures (84 dB SPL) can hide massive degeneration of cochlear nerve fibers, and that this neuropathy is exacerbated by loss of OC innervation (Usubuchi et al., ARO 2012). Here, we ask whether OC feedback also minimizes neural degeneration in mice aged without any purposeful acoustic overexposure.

Methods
In one group of 6-wk old CBA/CaJ mice, OC efferents were lesioned unilaterally via stereotaxic injections of melittin into the LSO and/or the VNTB, where the lateral and/or medial OC fibers originate, respectively. In another group, the crossed OC bundle was cut via midline section at the floor of the IVth ventricle. A third (control) group had no brainstem surgery. ABRs and DPOAEs were measured in both ears at 8, 11, 16, 22, 32 and 45 wks of age. The strength of medial OC efferents was assessed by shocking the OC bundle while measuring DPOAE amplitudes. Cochlear nerve degeneration, and the degree of de-efferentation, were assessed by immunostaining cochlear whole mounts for pre-synaptic ribbons (CtBP2), post-synaptic AMPA-receptor patches (GluR2) and OC terminals (VAT).

Results
At 45 wks, controls ears showed <15% loss of afferent synapses throughout the cochlea [8-32 kHz] and minimal (<8 dB) threshold shifts. In mice with partial or complete de-efferentation, threshold shifts were larger (10-30 dB) especially at the lowest and highest frequencies, and synaptic loss was significantly worse (by as much as 25%) especially in the apical and basal thirds of the cochlea. Analysis to date suggests that both MOC and LOC systems contribute to this effect.

Conclusion
These findings suggest that OC feedback may be essential for the long-term survival of cochlear nerve terminals in the face of a constant excitotoxic challenge from their highly active glutamatergic synapses with inner hair cells.

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Using the Auditory Steady-State Response to Measure Noise-Induced Auditory Nerve Degeneration
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Background
Moderate noise exposure can cause permanent loss of auditory nerve (AN) fibers, without causing hair cell loss or permanent threshold shifts (Kujawa & Liberman, JNeurosci 29:14077, 2009). This neuropathy is likely present in humans, but undetected by routine clinical examination. Reduction of suprathreshold auditory brainstem response (ABR) wave I amplitude has been used to quantify this primary neural degeneration in mice. However, ABR amplitudes are more variable in humans, and therefore may not be a useful diagnostic tool. This noise-induced neuropathy is selective for high-threshold ANs, i.e. those with low spontaneous rates (SRs) (Lin et al., ARO 2011, #400). Since low-SR fibers show greater phase locking to amplitude modulation than high-SR fibers (Joris & Yin, JAcoust Soc Am, 91:215, 1992), we hypothesized that the auditory steady-state response (ASSR), the gross electrical potential in response to amplitude-modulated tones, may be a useful metric of noise-induced AN loss.

Methods
8-wk mice were exposed to 98 dB SPL, 8-16 kHz noise for 2 hours. ASSRs and ABRs were recorded before, and 2 weeks after, exposure. ABRs were measured at tone pip frequencies of 11.3 and 32 kHz. ASSRs were evoked with carrier frequencies of either 11.3 or 32 kHz. Modulation frequencies were greater than 700 Hz so that ASSRs would be dominated by more peripheral auditory structures. Cochlear neuropathy was assessed by confocal microscopy of whole mounts immunostained for pre- and post-synaptic markers.

Results
This noise exposure causes loss of up to 50% of AN synapses with inner hair cells throughout the basal turn, i.e. at cochlear frequencies from 16 – 45 kHz. While ABR and ASSR amplitudes in response to 11 kHz tone pips or carriers, respectively, were similar to controls, at 32 kHz both ABR and ASSR amplitudes were significantly lower in noise-exposed mice. The phase consistency of the ASSR response at 32 kHz was also significantly lower in noise-exposed mice. To simulate the amplitude variability in humans, we randomly attenuated the responses, while holding the noise level constant. This did not affect the group difference in ASSR phase consistency, but eliminated the group difference in ASSR and ABR amplitudes.

Conclusion
Loss of ANs reduces ABR amplitude, ASSR amplitude, and ASSR phase consistency. Only ASSR phase consistency is unaffected by amplitude variability, and therefore might be used to diagnose AN loss in humans. Supported by NIDCD R01DC002258, R01DC000188, & P30DC005209.

Neurotrophin-Like Rescue of Spiral Ganglion Neurons after Noise
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Background
Neurotrophins can delay or prevent cochlear nerve degeneration secondary to hair cell loss and encourage dendritic sprouting in a denuded cochlear epithelium (Wise et al 2005). This raises the exciting possibility that, for noise exposures that damage neurons without destroying hair cells (Kujawa and Liberman 2009), delivery of neurotrophins or like-acting compounds could lead to hair cell reinnervation and functional recovery of IHC-afferent communications. Amitriptyline is a compound that mimics the neuroprotective effects of neurotrophins by Trk receptor agonist activity and has better bioavailability when delivered to the systemic circulation. Systemic amitriptyline suppresses neuronal apoptosis in a TrkA-dependent manner in hippocampus after glutamate agonist-induced neuro-excitotoxicity (Jang et al 2009). Here, we test whether acute and chronic neuro-excitotoxic consequences of noise exposure can be similarly suppressed by systemic amitriptyline.

Methods
Experiments were conducted on male, 16 wk old CBA/CaJ mice. Animals received once daily i.p. injections of amitriptyline (15 mg/kg) or an equal volume of vehicle (saline) for 5 days. Awake animals were noise exposed (8-16 kHz OBN, 100 dB SPL, 2 h) on day 2 and were held for various post-exposure times (1 d to 1 yr) before physiologic testing (ABR, DPOAE). Tissues were retrieved and studied as immunostained (CtBP2, Na-K-ATPase) cochlear whole mounts.
and osmium-stained, plastic-embedded sections (10 µm). We counted hair cells, synaptic ribbons and spiral ganglion cells at cochlear places corresponding to 5.6, 11.3 and 32 kHz frequencies.

Results
Acute noise-induced threshold shifts were similar in amitriptyline- and saline-treated animals, they recovered to baseline by 2 wk post exposure and changed similarly with age throughout our period of monitoring. As expected, hair cells were not lost from this exposure and numbers were not altered by drug treatment. One year after noise, ribbon counts were ~20-40% greater and ganglion cell counts were ~25-40% greater, depending on frequency, in drug-treated ears vs. saline-treated controls. Although nerve survival appears enhanced in amitriptyline-treated ears, a corresponding functional preservation of suprathreshold ABR amplitudes was not observed using this treatment regimen.

Conclusion
Treatment with amitriptyline, which has neurotrophin-like activity at TrkA receptors, preserves spiral ganglion neurons in noise-exposed mice. Studies aimed at optimizing the dosing regimen and extending characterization of the functional status of these neurons are underway. In parallel, a cochlear explant preparation is in use to characterize drug effects on dendritic sprouting and afferent connections. Research supported by NIDCD: R01 DC008577

Glutamate-induced neural degenerations in cochlear organotypic cultures
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Background
Glutamate is believed to be the excitatory amino acid neurotransmitters released by inner hair cells. When glutamate is released, it can bind to 2 classes of glutamate receptors (AMPA or Kainate) that are located on auditory nerve fiber synapses beneath the inner hair cell. During intense sound stimulation, excess glutamate release can lead to excitotoxicity. Excitotoxicity can be induced in vivo by infusing glutamic acid or kainic acid into the cochlea; this results in significant swelling at type I afferent synapses or death of type I spiral ganglion neurons. Little is currently known about glutamate and kainate excitotoxicity in developing afferent synapses of the postnatal ear. To address this question, we treated postnatal cochlear organotypic cultures with glutamic or kainic acid and determined their effects on the afferent terminals and spiral ganglion neurons (SGN).

Methods
Cochlear explants from postnatal day 3 rats were treated with glutamic acid (10-5000 µM) or kainic acid (100-20000 µM) for 24 h. Cochlear hair cells, auditory synapses, and spiral ganglion neurons (type I and type II) were labeled and evaluated by confocal microscopy.

Results
Treatment of rat cochlear cultures with glutamic acid resulted in damage that was confined to auditory nerve synapses and peripheral nerve fibers. In contrast, cochlear hair cells and SGN soma appeared normal. Surprisingly, kainic acid failed to induce obvious damage to hair cells, synapses, or SGN soma.

Conclusion
The lack of kainic acid-induced damage suggests that there is a paucity of functional kainate receptors on the afferent synapses and soma of SGN. Alternatively, glutamic acid only damaged the synaptic terminal of auditory nerve fibers. These results suggest that functional AMPA-type glutamate receptors are present mainly at the afferent synapse.
Involvement of the Superoxide Radical in Salicylate-Induced Spiral Ganglion Degeneration

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Background
Salicylate, the active component in aspirin, is a potent antioxidant that can protect against cochlear hearing loss. However, high doses of salicylate have long been known to cause transient tinnitus and hearing loss. Recent studies have shown that salicylate does not damage cochlear hair cells or supporting cells, but paradoxically selectively destroys spiral ganglion neurons (SGN) by mechanisms consistent with apoptosis. To gain insights into the processes that initiate SGN death, we monitored superoxide generation and characterized the early stages of caspase-mediated SGN death.

Methods
Cochlear organ cultures were prepared from postnatal day 3 rats and treated with different concentrations of salicylate. Superoxide, a reactive oxygen species, was detected with dihydroethidium, a fluorescent probe. Fluorengenic caspase-specific probes were used to detect initiator caspase 8 and caspase 9 and downstream executioner caspase 3.

Results
Salicylate caused a dose and time-dependent loss of SGN and nerve fibers. SGN degeneration was characterized by nerve fiber disintegration and nuclear shrinkage and fragmentation. Caspase 8 and caspase 9 were activated during the early stages of degeneration followed by activation of executioner caspase 3. Salicylate significantly increased superoxide labeling in SGN. MnTMPyP pentachloride, a cell-permeable scavenger of superoxide, significantly attenuated salicylate-induced SGN degeneration.

Conclusion
One of the most puzzling aspects of salicylate ototoxicity is its selective damage of SGN. Our results suggest that SGN are preferentially damaged because of the increase of superoxide exclusively in SGN. This interpretation is supported by the fact that MnTMPyp, a superoxide scavenger, prevented salicylate-induced SGN degeneration.

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Exogenous BDNF and NT-3 have Distinct Biological Effects on Afferent Synaptogenesis on Inner Hair Cells (IHCs) without Endogenous NT-3 in vitro

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Background
Previous results (Wang & Green, J. Neurosci. 31: 7938-49, 2011) using neonatal rat organotypic cochlea explant cultures indicated that endogenous NT-3 in the postnatal organ of Corti has a distinctive role, not mimicked by BDNF, in promoting SGN synapse regeneration on IHCs. Here we ask directly whether BDNF and NT-3 are equivalent in their ability to promote SGN axon growth and synaptogenesis on IHCs.

Methods
We generated Atoh1-CreERT²; NT-3³/², Z/EG mice and treated with tamoxifen on P0 to delete NT-3 from a random subset of cochlear IHCs and label the NT-3-lacking hair cells with GFP. Organotypic cochlear explant cultures were made from the mice three days after tamoxifen injection. IHC-SGN synapses were destroyed after 1 day in vitro (DIV) with the glutamate agonist kainate (0.5 mM, 30 min). Reinnervation of IHCs was assessed after 3 more DIV with or without presence of neurotrophins (NT-3 or BDNF). SGN fiber-IHC contacts and postsynaptic densities (PSDs) on individual IHCs were quantified in confocal image stacks. These counts were compared between NT-3-lacking IHCs and adjacent IHCs expressing NT-3.

Results
Although reinnervation after excitotoxic trauma is generally poor in mice, most NT-3-lacking IHCs were selectively bypassed by regrowing SGN axons, while adjacent NT-3-expressing IHCs were reinnervated and acquired new PSDs. Significantly (p<0.05) fewer PSDs (0.06 ± 0.3, n=87) were associated with reinnervated IHCs lacking NT-3. Culturing with NT-3 enhanced both SGN fiber regrowth (1.4±1.2 SGN fibers/IHC, n=111) and synaptogenesis (1.3±1.5 PSDs/ IHC, n=64) on NT-3-lacking IHCs to a level comparable to NT-3-expressing IHCs (1.2±1.1 SGN fibers/IHC, n=499; 0.9±1.4
PSDs/ IHC, n=342). In contrast, culturing with BDNF had no significant effect on synaptogenesis on NT-3-lacking IHCs (0 PSDs/IHC, n=30) although BDNF did promote SGN fiber growth to a similar extent as did NT-3.

Conclusion
Endogenous NT-3 expressed in each IHC promotes SGN synapse regeneration on that IHC after synapse destruction by excitotoxic trauma. Addition of exogenous NT-3 or BDNF to the cultures during the three days over which reinnervation occurs results in comparably improved SGN fiber growth. However, only NT-3 addition, but not BDNF addition, enhances synaptogenesis on IHCs.

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Gene Expression Changes in the Spiral Ganglion (SG) After Deafening
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Background
Hair cells (HCs) are the sole afferent input to spiral ganglion neurons (SGNs). Following aminoglycoside exposure from postnatal day 8 (P8) to P16, SGNs degenerate and gradually die. In rats, this occurs over a period of ~3 months (Alam et al. JCN 503: 832-52, 2007). SGN death may be due to loss of HC-derived neurotrophic factors (NTFs). However, expression of NTFs other than NT-3 persists throughout the period during which most SGNs die. Possibly, SGN death is an indirect outcome of HC loss, due to degenerative changes in the cochlea initiated by HC loss. We used microarray-based gene expression profiling to identify changes in the SG during the SGN death period and gain insight into why SGNs die.

Methods
Rats were deafened as previously described (Alam et al., 2007) and euthanized at P32, when SGN loss has just become significant or at P60, a time at which ~50% of the SGNs remain. Spiral ganglia were removed and divided into apical and basal halves prior to extraction of RNA. cDNA was hybridized in triplicate to Nimblegen HD2 rat gene expression arrays. Three separate biological repetitions were performed. DNASTAR software was used for quantile normalization and DAVID for functional annotation.

Results
13,000-15,000 genes are expressed above background. In comparing expression changes between P32 and P60 (maturation-related) or between hearing and deaf, expression of most genes does not change significantly. This includes genes encoding most NTFs and their receptors and genes related to apoptosis/survival, implying that lack of NTFs is not a primary cause of SGN death. Expression of ~8% changes significantly in hearing rats between P32 and P60, implying significant changes due to maturation. Expression of ~5% changes significantly after deafening at P32; remarkably, most transcriptional changes occurring after deafening also occur in maturation, implying accelerated maturation-related changes in the SG post-deafening. <3% of expression changes are deafness-specific. Many of these are indicative of infiltration of immune system-related cells into the spiral ganglion.

Conclusion
Gene expression profiling has provided significant insight into the question of why SGNs die after HCs are killed by aminoglycosides. The data do not support a hypothesis that SGNs die because of loss of HC-derived NTFs in spite of observations that NTF application can rescue SGNs. Rather, SGN death may be due more directly to changes in glia or other cell types in the deafened ganglion.

1026
Increased presence of cells of the immune system in the spiral ganglion during spiral ganglion neuron (SGN) death post-deafening
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Background
Following hair cell (HC) death due to aminoglycoside exposure from postnatal day 8 (P8) to P16, rat SGNs degenerate and die over a period of ~3 months (Alam et al. JCN 503: 832-52, 2007). The reason for SGN death is not clear. Several neurotrophic factors remain expressed in the cochlea after HC loss and SGNs do not die in all types of HC death. Possibly, SGN death is an indirect outcome of HC loss, due to other degenerative changes in the cochlea. We used
microarray-based gene expression profiling to identify changes in the spiral ganglion during the SGN death period and gain insight into why SGNs die.

**Methods**
Rats were deafened by daily kanamycin injection P8-P16 and euthanized at P32, when SGN loss is just becoming significant, or at P60, when ~50% of the SGNs have died. RNA was isolated from dissected spiral ganglia. cDNA was hybridized in triplicate to Nimblegen HD2 rat gene expression arrays with three separate biological repetitions. DNASTAR software was used for quantile normalization and DAVID for functional annotation. Expression of selected representative genes was verified by qPCR. Immunofluorescence was performed as previously described (Alam et al., 2007).

**Results**
For genes expressed in the spiral ganglion, expression of ~0.8% decreases by >2X and of ~2% increases by >2X in deafened P60 rats, relative to hearing littermates. Many of these are indicative of infiltration of immune system-related cells into the spiral ganglion. For example, C-C and C-X-C chemokines are upregulated (CCL3 verified by qPCR) and interferon-inducible genes, e.g., Mx1, Mx2, are upregulated. There is a resident population of macrophages in the ganglion and characteristic gene expression (e.g., IBA1/AIF1, CD14, CD11b/ITGAM, CD45/PTPRC, CD68) is evident in the gene expression profiling data in hearing rats. Expression of several of these, including PTPRC and CD68 (verified by qPCR), as well as MHCII expression increases >2X by P60 after deafening, suggesting an increased number of activated macrophages. This was verified by detection, by immunofluorescence, of an increased number of CD68-positive cells in the deafened spiral ganglion at P60.

**Conclusion**
Gene expression profiling provides evidence for increased immune cell presence, including cells of the innate system, in the spiral ganglion at the time when SGNs are dying. While their presence may be related only to clearance of debris, the observations raise the possibility that they are involved in SGN death.
Targeted deletion of oncomodulin alters efferent innervation in the cochlea
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Background
The tight regulation of Ca\textsuperscript{2+} is essential for cochlear function, and yet the role of Ca\textsuperscript{2+} binding proteins in hair cells remains elusive. Oncomodulin (Ocm), a member of the parvalbumin family, has a restricted expression pattern, and in the cochlea is found in the basolateral membrane and hair bundle of outer hair cells (OHCs). Oncomodulin may mediate efferent-induced OHC responses and other Ca\textsuperscript{2+} dependent mechanisms, and therefore in the absence of Ocm, OHC function and efferent-mediated responses may be disrupted. As an initial test of this hypothesis, we performed preliminary morphological studies on OHCs and their efferent innervation in both wild-type and Ocm\textsuperscript{-/-} mice.

Methods
To study the role of Ocm, we generated a conditional Cre-lox knockout line with Cre-recombinase driven by the β-actin promoter (Ocm\textsuperscript{tm1.Ddsi}). The absence of Ocm expression was confirmed by RT-PCR and immunocytochemistry. Hearing function was assessed by auditory brainstem responses (ABRs). Choline acetyltransferase (ChAT) was used to label efferent axons and terminals. Phalloidin labeling was used to assess hair cell loss.

Results
Preliminary analysis of Ocm\textsuperscript{tm1.Ddsi} mice reveals no obvious phenotypic abnormalities compared with controls. At 4 weeks, there is no difference in cochlear function between mutant and wild-type littermates. However by 8 - 10 weeks, Ocm\textsuperscript{-/-} mutants have ABR threshold shifts of as much as 60 dB in frequencies above 10 kHz. There is no significant hair cell loss prior to 8 weeks. However after 8 weeks, OHC loss is pronounced. Ocm\textsuperscript{-/-} mutants exhibit abnormal ChAT-labeled, efferent innervation patterns compared to controls. At 4 weeks, Ocm\textsuperscript{-/-} mutants also exhibit a dense innervation in the IHC region and a more disorderly innervation below OHCs. We see a higher density of efferent terminals below OHCs at 1 month compared to 4 months and in the apex compared to the base. The number of terminal clusters in basal regions begins to decline precipitously around 2 months. Also efferent terminal clusters in the Ocm mutant are smaller as the animal ages. Finally, there is significant remodeling of efferent projections in the cochlea in the 3 and 4 month Ocm mutants.

Conclusion
In the absence of a major Ca\textsuperscript{2+} binding protein, efferent organization is disrupted and OHCs are more susceptible to damage. Our data suggest that calcium regulators such as Ocm may play a protective role in hearing and play a role in organizing efferent innervation.
Human Spiral Ganglion Degeneration is Independent of Hair Cell Loss
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Background
Human spiral ganglion cells, in contrast to those of experimentally deafened animals, do not degenerate shortly after the loss of the organ of Corti. In fact, regardless of the time interval following the loss of hair cells and the individual’s death, there is not an associated degeneration of spiral ganglion neurons (SGN). Numerous investigations with animal models have shown that cochlear neurons degenerate following degeneration of hair cells induced by ototoxicity. The time interval from hair cell loss to neuron loss in animals varies from 25% of the animal’s life expectancy in the guinea pig to 1% in the chinchilla. In contrast to animals, we have observed surviving SGN in human temporal bone specimens from congenitally deaf humans with no surviving hair cells.

Methods
Histopathological and clinical history investigation of 40 temporal bones in our laboratory that are from patients, deafened by various etiologies, with no surviving hair cells. The durations of the hearing losses ranged from 9 to 93 years.

Results
Ganglion cell counts varied from 360 to 28,262 (avg. 14,514; SD=7934.75); the normal range for humans is 13,918 to 36,918, depending on age. Duration of hearing loss ranged from 5 to 93 years with an average of 38 years (SD=22.8), and hearing thresholds were recorded for durations ranging from 2 to 39 years (avg. 15.6 years; SD=10.6). Statistical analysis demonstrated no relationship between SGN survival and years of hearing loss.

Conclusion
Eight temporal bones from individuals that were deaf for an average of 21.25 years had SGN counts within the normal range for their age. SGN survival ranging in absolute number from 11,722 to 19,093 was demonstrated in congenitally deaf individuals who survived 89-93 years of age with no surviving hair cells. These numbers suggest that, unlike animals, human SGN do not undergo degeneration following loss of auditory hair cells. Further investigations are necessary to determine factors involved in human SGN survival after hair cell degeneration.
Cochlear Uptake of Gentamicin in Neonatal Mice
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Background
Unlike the functionally mature cochlea of human newborns, the murine cochlea becomes functionally mature around 2 weeks after birth. In mice, tight junctions between adjacent marginal cells, and between adjacent endothelial cells, within the stria vascularis are established during this postnatal period, with consequent enrichment of endolymph with potassium. This dynamic postnatal cochlear maturation in the mouse provides a unique opportunity to study how ototoxic reagents permeate the developing blood-labyrinth barrier to reach young sensory hair cells.

Methods
Neonatal C57Bl/6 mice were intraperitoneally injected with fluorescently-conjugated gentamicin (GTTR) for 30 or 60 minutes at various postnatal ages prior to in situ fixation. Harvested cochlear tissues were either whole-mounted or cryosectioned for confocal microscopy. The intensity and distribution of GTTR in all tissues were acquired at the same confocal settings in all experimental groups. The intensity of GTTR fluorescence in cochlear and renal proximal tubule tissues were quantified and compared between different age groups (p7, p21, p35, n=6, 3, 6 respectively).

Results
GTTR fluorescence in hair cells and the lateral wall (including marginal, intermediate and basal cells of the stria vascularis, and fibrocytes of the spiral ligament) was significantly higher at p7 than at p21 or p35 (p<0.01). This is suggestive of paracellular permeation of GTTR through the premature blood-labyrinth barrier. Furthermore, hair cells exhibited a heterogeneous pattern of GTTR uptake between p7 to p10. That is, a subset of sensory and supporting epithelial cells presented increased GTTR fluorescence compared to adjacent cells of the same cell type. Animals treated with Texas Red only had negligible red fluorescence at all time points. Renal proximal tubules did not show significant differences in fluorescence intensity between different time points.

Conclusion
Prior to complete maturation of the cochlear blood-labyrinth barrier, GTTR readily permeates into strial cells, fibrocytes and sensory hair cells. A heterogeneous pattern of GTTR uptake in the organ of Corti from p7 to p10 corresponds with a known wave of purinergic activation of sensory epithelial cells that precedes the onset of functional hearing. This suggests that purinergic ion channels, such as P2X2, may facilitate hair cell uptake of GTTR and aminoglycosides.

Identification of Cisplatin-Binding Proteins in the Cochlea
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Background
Cisplatin is widely used as an antineoplastic drug, but its ototoxic and nephrotoxic side-effects, as well as the inherent or acquired resistance of some cancers to cisplatin, remain significant clinical problems. Cisplatin’s selectivity in killing rapidly proliferating cancer cells is largely dependent on covalent binding to DNA via cisplatin’s chloride sites that had been aquated. We hypothesized that cisplatin’s toxicity in slowly proliferating or terminally differentiated cells is primarily due to drug-protein interactions, instead of drug-DNA binding.

Methods
To identify proteins that bind to cisplatin, we synthesized two different platinum-agarose conjugates, one with two amino groups and another with two chlorides attached to platinum that are available for protein binding, and conducted pull-down assays using cochlear and kidney cells, including the organ of Corti HEI-OC1 cells. The proteins that bound to the platinum-agarose were removed from platinum-agarose in SDS sample buffer, analyzed by SDS-gel electrophoresis and Coomassie blue staining, and identified by mass spectrometry.

Results
Platinum-agarose pull-down assay revealed that proteins poorly bound to cisplatin at the chloride site, while many proteins bound to the amino groups with high affinities. Comparison among different cell lines did not show significant
differences in binding proteins, except for myosin IIA, which only appeared in HEI-OC1 cells. Mass spectrometric analysis on protein bands after gel electrophoresis and Coomassie blue staining identified several proteins, including myosin IIA, glucose-regulated protein 94 (GRP94), heat shock protein 90 (HSP90), calreticulin, valosin containing protein (VCP), and ribosomal protein L5, as cisplatin-binding proteins in HEI-OC1 cells. Previous reports suggested that some of these proteins are specifically expressed in the cochlea and most of them are overexpressed in cancer.

Conclusion
We have identified several cisplatin-binding proteins that are either specifically expressed in the cochlea or overexpressed in cancer. These proteins likely play a role in cisplatin-induced ototoxicity/nephrotoxicity or cisplatin resistance of cancer. Future studies on the cisplatin-binding proteins will elucidate whether these drug-protein interactions are involved in ototoxicity and nephrotoxicity, or contribute to tumor resistance to cisplatin treatment.

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[1031]
Early time course of gentamicin uptake in peripheral vestibular cells after systemic administration
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Background
Systemic aminoglycoside therapy can induce vestibulotoxicity as well as cochleotoxicity, resulting in imbalance and visual dysfunction. The underlying trafficking routes and cytotoxicity mechanisms remain unknown. We investigated the early time course of gentamicin trafficking into the peripheral vestibular system after systemic administration of fluorescently-tagged gentamicin (GTTR).

Methods
C57BL/6 mice received GTTR 2 mg/kg via intraperitoneal injection; control mice received equivalent doses of Texas Red or PBS. Mice were sacrificed at 0.5, 1, 2, 3 or 4 hours, followed by transcardiac perfusion with PBS, and fixation with 4% paraformaldehyde. Excised vestibular tissues were permeabilized and subsequently counterlabeled with fluorescent phalloidin, and examined by sequential scanning laser confocal microscopy. Identical confocal settings were used for all experimental and control groups. Fluorescence pixel intensities in nonsensory and sensory cells (including dark, transitional, and supporting cells) from single optical sections were obtained after removal of extracellular pixels. Mean pixel intensity-time course plots were constructed.

Results
Vestibular cells of both ears of all animals exposed to GTTR exhibited changes in GTTR fluorescence over time. Low intensity fluorescence was detected at 0.5 hours, and increased in intensity to peak at 3 hours, before declining 4 hours after injection. The distribution of GTTR fluorescence differed in sensory and nonsensory cells. Sensory cells typically had only diffuse GTTR fluorescence, at all tested timepoints. In contrast, nonsensory cells displayed (i) an intensely fluorescent granular distribution (> 0.6 microns in diameter), as well as (ii) a diffuse pattern of fluorescence throughout the cytosol. The area of granular fluorescence in dark cells and transitional cells significantly increased over time. Control vestibular tissues exposed to PBS or hydrolyzed Texas Red had negligible fluorescence at all tested timepoints.

Conclusion
Systemically-administered GTTR is rapidly taken up by murine peripheral vestibular sensory and nonsensory cells, peaking in fluorescence intensity 3 hours after a single injection. The intensity of GTTR in sensory and nonsensory cells share similar time courses. However, the distribution of GTTR within cells is very different. Diffuse fluorescence suggests entry via GTTR-permeant ion channels, while granular fluorescence suggest either endocytosis and/or compartmentalization of cytosolic GTTR. The difference in GTTR distribution within sensory and nonsensory cells suggests differential uptake mechanisms and/or subcellular GTTR targets in these cell types. The intra-vestibular trafficking routes of GTTR also remain to be elucidated.

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Is a Temporary Threshold Shift Required for Sound-Enhanced Uptake of Gentamicin?
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Background
Animal studies have firmly demonstrated the ototoxic synergy between acoustic overstimulation and aminoglycoside antibiotics. We have previously reported that prior sound exposure, at moderate sound levels which induce temporary threshold shifts (TTS), also enhanced gentamicin uptake in the cochlea. We propose that sound-enhanced aminoglycoside uptake by the cochlear tissue is a candidate mechanism for ototoxic synergy. In order to determine whether correlation between TTS and sound-enhanced gentamicin uptake is present, we systemically varied the sound level of wideband noise, and examined its effect on cochlear uptake of gentamicin.

Methods
To induce temporary threshold shifts, adult C57Bl/6 mice were exposed to wideband noise (86–96 dB SPL) for 6 hours/day for 3 days. The degree of TTS was assessed by tonal ABR measurements (4, 8, 12, 16, 24 and 32 kHz) at multiple time points before and after sound exposure. Mice were also intraperitoneally injected with fluorescently-conjugated gentamicin (GTTR) after sound exposure. Thirty minutes later, fixed cochlear tissues were excised and processed for confocal microscopy. The intensity of GTTR fluorescence were acquired using identical confocal settings, quantified and statistically tested to determine the effect of sound exposure on GTTR uptake in the cochlea.

Results
Higher sound levels (91 and 96 dB SPL) produced evident TTS at higher (16, 24 and 32 kHz) frequencies, while no TTS was observed at lower frequencies (4 and 8 kHz). Sound levels that induced TTS also enhanced hair cell uptake of GTTR. For cochleae with TTS, we examined multiple frequency locations along the organ of Corti. Enhanced hair cell uptake of GTTR was observed in basal high frequency regions with TTS, and also in more apical low frequency regions with normal ABRs thresholds. In cochleae without evident TTS, hair cells did not show the enhanced uptake of GTTR that is associated with TTS.

Conclusion
TTS is required for sound-enhanced uptake of gentamicin by hair cells in the cochlea. Sound-enhanced uptake of gentamicin by hair cells is not specific to the frequency regions with TTS, and also occurs at frequencies below the region of TTS. Thus, the mechanisms that induce sound-enhanced gentamicin uptake extend beyond the cochlear region associated with TTS and the mechanisms that induce TTS.

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Bumetanide hyperpolarizes cells and enhances cellular uptake of gentamicin by elevating cytosolic Ca²⁺ and facilitates intermediate conductance Ca²⁺-activated potassium channels
Tian Wang¹, Yu-qin Yang¹, Takatoshi Karasawa¹, Meiyan Jiang¹, Peter Steyger¹, Zhi-Gen Jiang¹
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Background
Loop diuretics such as bumetanide and furosemide enhance aminoglycoside ototoxicity when co-administered to patients and animal models. The underlying mechanism(s) is poorly understood.

Methods
We investigated the effect of these diuretics on cellular uptake of aminoglycosides, using fluorescently-tagged gentamicin (GTTR), and the action of these diuretics on membrane potentials and specific channel currents using intracellular/whole-cell recordings of Madin-Darby Canine kidney (MDCK) cells. MDCK cells preloaded with the calcium indicator Fluo-4 were used to detect the effect of bumetanide on cytosolic Ca²⁺-level by image recording with a Yokogawa CSU10 spinning disk confocal system, and data quantitation and analysis with Fiji/ImageJ software.

Results
We found that bumetanide and furosemide concentration-dependently enhanced cytoplasmic GTTR fluorescence up to ~60%. This enhancement was suppressed by La³⁺, a non-selective cation channel (NSCC) blocker, and by K⁺ channel blockers Ba²⁺ and clotrimazole, but not by tetraethylammonium (TEA), 4-aminopyridine (4-AP) or glipizide, nor by Cl⁻ channel blockers diphenylamine-2-carboxylic acid (DPC), niflumic acid (NFA), and CFTRinh-172. Bumetanide and furosemide hyperpolarized MDCK cells by ~14 mV, increased whole-cell I/V slope conductance; the bumetanide-induced
net current I/V showed a reversal potential \( V_r \) \( \sim -80 \text{ mV} \). Bumetanide-induced hyperpolarization and I/V change was suppressed by \( \text{Ba}^{2+} \) or clotrimazole, and absent in elevated \([\text{Ca}^{2+}]_i\) (2 \( \mu \text{M}\)), but not affected by apamin, 4-AP, TEA, glipizide, DPC, NFA or CFTR \text{inh}-172. Bumetanide and furosemide stimulated a surge of Fluo-4-indicated cytosolic \( \text{Ca}^{2+} \). \( \text{Ba}^{2+} \) and clotrimazole alone depolarized cells by \( \sim 18 \text{ mV} \) and reduced I/V slope with a net current \( V_r \) near -85 mV, and reduced GTTR uptake by \( \sim 20\% \). \( \text{La}^{3+} \) alone hyperpolarized the cells by \( \sim 14 \text{ mV} \), reduced the I/V slope with a net current \( V_r \) near -10 mV, and inhibited GTTR uptake by \( \sim 50\% \). In the presence of \( \text{La}^{3+} \), bumetanide caused negligible potential or I/V change.

**Conclusion**

We conclude that NSCCs constitute a major cell entry pathway for cationic aminoglycosides; bumetanide enhances aminoglycoside uptake by hyperpolarizing cells that increases cation influx driving force. Bumetanide-induced hyperpolarization is caused by elevating intracellular \( \text{Ca}^{2+} \) and thus enhances intermediate conductance \( \text{Ca}^{2+} \)-activated K\(^+\) channels.

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**Protection of Hair Cells from Aminoglycoside Antibiotics in the Zebrafish merovingian Mutant**

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**Background**

A number of therapeutic agents, including aminoglycoside antibiotics and chemotherapeutics cause hair cell death. While progress has been made in delineating the cellular pathways responsible for toxicant induced hair cell death, our knowledge is still incomplete, and hearing loss remains a dose limiting side effect of multiple therapeutic drugs. Our lab uses the zebrafish lateral line system to identify novel genes involved in toxicant induced hair cell death.

**Methods**

5-6 days post fertilization zebrafish are used for all experiments. For experiments looking at neomycin toxicity, fish were exposed to varying doses of neomycin for 30 minutes, and then given 1 hour to recover. Hair cells were labeled using parvalbumin staining to quantify hair cell number after treatment. For measuring uptake of Texas Red-conjugated neomycin or FM1-43 fish were exposed to the compound for 15 minutes or 1 minute respectively and then imaged immediately.

**Results**

We have previously identified the *merovingian* (*mero*) mutant in a screen for zebrafish mutants where lateral line hair cells are protected from aminoglycoside hair cell death (Owens et al, PLOS Genetics, 2008). *merovingian* mutants show complete protection against neomycin-induced hair cell death which appears to be due to a failure of the drug to enter hair cells. These mutants also show a decreased initial number of lateral line hair cells, vestibular defects, and decreased FM1-43 uptake suggesting impaired hair cell development and function. Through genetic mapping we have identified a mutation in transcription factor *gcm2*, a gene necessary for global pH and ion regulation in the *merovingian* mutants. This is of particular interest as we have previously identified a mutation in the chloride/bicarbonate exchanger, SLC4A1b, as also conferring resistance to aminoglycoside induced hair cell death (Hailey et al., PLOS Genetics 2012, Owens et al, PLOS Genetics, 2008).

**Conclusion**

Our results implicate a role for global ion and pH regulation in controlling hair cell function as well as susceptibility to aminoglycoside-induced hair cell death. We are continuing to study the *merovingian* mutant to elucidate the relevant ionic transporters affecting hair cell function and aminoglycoside susceptibility.
Neutral sphingomyelinase is involved in hair cell death induced by gentamicin
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**Background**
The sphingolipid metabolites ceramide, sphingosine, and sphingosine-1-phosphate (S1P) are known as a new class of lipid second messengers and reportedly play essential roles in the regulation of cell proliferation, survival, and death. Ceramide has been shown to regulate diverse cellular processes including apoptosis, cell senescence, the cell cycle, and cellular differentiation. Recently, we showed that exogenous ceramide increased hair cell death in gentamicin ototoxicity (Nishimura et al., 2010). Sphingomyelinase (SMase) converts sphingomyelin to ceramide. Several signaling molecules, such as tumor necrosis factor, Fas ligand, γ-interferon, interleukin 1, vitamin D, and stressful events including radiation and ischemia, reportedly activate SMase and promote ceramide-induced apoptosis. This study was designed to investigate the possible involvement of neutral SMase in hair cell death due to gentamicin.

**Methods**
The basal turn of the organ of Corti was dissected from Sprague–Dawley rats on postnatal days 3 (p3) to 5 (p5). Cochlear cultures were exposed to a medium containing 35 μM gentamicin for 48 hours to assess the effects of GW4869, a neutral SMase inhibitor.

**Results**
One to 30 μM GW4869 did not affect hair cell loss induced by gentamicin. However, hair cell loss significantly decreased in the presence of 35 to 50 μM GW4869.

**Conclusion**
The present finding strongly suggests that GW4869 protected hair cells against gentamicin ototoxicity by inhibiting ceramide synthesis via the SMase pathway.

Targeted activation of ERK1/2 in supporting cells is sufficient to promote hair cell death in the adult utricle
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**Background**
In the neonatal cochlea, extracellular regulated kinases 1 and 2 (ERK1/2) are activated in supporting cells surrounding an epithelial wound, and also around hair cells damaged by the aminoglycoside, neomycin. Inhibition of MEK1/2, the upstream kinase, prevents activation of ERK1/2 and affords some protection of inner hair cells (Lahne & Gale 2008). Given that supporting cells exhibited the phosphorylated-ERK1/2 signal, we concluded that supporting cells could be contributing to hair cell death. Direct evidence that supporting cells can phagocytose hair cells was obtained using live-imaging of ototoxically damaged avian utricles (Bird et al. 2010). Here we investigate whether a similar supporting cell and ERK1/2-dependent mechanism is present in the mature inner ear and whether activating ERK in supporting cells is sufficient to trigger phagocytic activity.

**Methods**
Adult and neonatal mouse utricle explant cultures were used to examine ERK1/2 activation following acute damage to hair cells using a multi-photon laser or using neomycin for 24 hours. The MEK1/2 inhibitor U0126 was used to probe the function of ERK1/2 activation. In order to determine the role of ERK1/2 in supporting cells, we used an adenoviral protocol (Brandon et al. 2012) to infect supporting cells with consecutively active MEK1 (CA-MEK1). Infected utricles were fixed and analysed using immunohistochemistry and confocal microscopy, or were subjected to live imaging using a spinning disk confocal microscope.

**Results**
Damaging hair cells using either the laser or neomycin resulted in the activation of ERK1/2 in the neighbouring supporting cells. Although we observed subtle differences in the spatiotemporal properties of ERK1/2-activation, the overall pattern was very similar in both mature and immature utricles. Pharmacological inhibition of MEK1/2 achieved a small but significant protection of hair cells during 24 hours aminoglycoside treatment in adult utricles. Constitutive activation of MEK1 in supporting cells using adenovirus was confirmed by phosphoERK staining. ERK1/2 activation resulted in
changes in cell phenotype including expansion of cell surfaces, movement towards the surface of the epithelium, and suggested phagocytic behavior. These phenotypes were confirmed in live imaging experiments. Finally, the activation of ERK1/2 in supporting cells resulted in a reduction in the number of hair cells in infected utricles.

Conclusion
This work provides evidence that supporting cells play a significant role in mediating hair cell death in adults, and that activation of the ERK1/2 signalling pathway is sufficient for this.
protein synthesis activity on a cell-by-cell basis. In combination with immunocytochemistry methods, BONCAT allows correlation of protein synthesis activity with markers of cellular signaling pathways.

Results
Using the BONCAT method, we discovered for the first time that AGs inhibit protein synthesis in hair cells. We found that protein synthesis inhibition is strictly correlated with the activation of the c-Jun kinase (JNK) pathway: shown by microscopy on a cell-by-cell basis, hair cells with arrested protein synthesis display strong JNK activation. In addition, we find that the ototoxic potency of a given aminoglycoside (tested were apramycin, kanamycin, gentamicin) correlated very well with its ability to cause cytosolic protein synthesis inhibition and JNK activation.

The observed stress response pattern (protein synthesis inhibition, JNK activation, caspase-3 mediated apoptosis) is characteristic for the so-called ribotoxic stress response, caused by a class of toxins that bind to and/or inactivate ribosomal RNA (rRNA). Indeed, AGs are known to bind to both mitochondrial and cytosolic rRNA. While the binding to mitochondrial rRNA has already been reported to contribute to AG ototoxicity, the affinity to cytosolic rRNA was believed to be too low to cause damage. Our data suggests the opposite, and we propose that a stress response similar to ribotoxic stress contributes to aminoglycoside ototoxicity. In agreement with this, we found that the FDA-approved anti-cancer drug sorafenib, known to inhibit ribotoxic stress response, also inhibits aminoglycoside-induced JNK activation and, to a limited degree, hair cell apoptosis.

Conclusion
In summary, we report the discovery of a novel stress pathway that contributes to aminoglycoside-induced hair cell degeneration. We demonstrate that aminoglycosides inhibit protein synthesis in hair cells and activate a stress pathway similar to ribotoxic stress response. A FDA-approved drug known to inhibit ribotoxic stress prevents JNK activation and improves hair cell survival.
Effects of dietary nutrients on amikacin ototoxicity in guinea pigs: reduced hearing loss and increased hair cell survival

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Background
Amikacin is an aminoglycoside antibiotic that is toxic to sensory cells in the inner ear, and produces hearing loss. Much of the damage to the sensory cells is the result of metabolic stress and related free radical production. Several agents that attenuate free radical production have been shown to attenuate amikacin and other aminoglycoside antibiotic induced ototoxicity. Here, we present data demonstrating protection from amikacin ototoxicity using a dietary supplement, composed of beta-carotene, vitamins C and E, and magnesium.

Methods
Albino guinea pigs were maintained on a nutritionally complete standard laboratory diet (2040 Teklad Global Guinea Pig Diet; n=10) or a custom-manufactured version of the 2040 diet (TD.08623) with increased levels of beta-carotene, vitamins C and E, and magnesium (n=8). Maintenance on the TD.08623 diet has previously been shown to reduce noise-induced and gentamicin-induced hearing loss in guinea pigs. Dietary manipulation was initiated 28-days prior to initiating daily amikacin injections. Subjects were maintained on the assigned diet during the daily amikacin injections (200 mg/kg/day amikacin x 28 days), and for the following 9 weeks after the last amikacin injection. Auditory brainstem response (ABR) thresholds were assessed at frequencies including 2, 4, 8, 16, and 24 kHz; tests were conducted at eight specific test times before, during, and after amikacin. At the end of the 9-week post-amikacin ABR assessment, subjects were euthanized, cochlear tissues were harvested, and hair cells were labeled and counted.

Results
Functional protection included a significant reduction in ABR threshold shift of ~10-20 dB across test frequencies and across test times. Inner hair cells (IHCs) were completely eliminated in amikacin-treated controls, whereas 80-90% were missing in animals receiving the supplemented diet plus amikacin. Outer hair cells (OHCs) were largely eliminated in the control group, except in the most apical 25% of the organ of Corti, where average OHC survival was ~10%. In animals receiving supplemented diet, average OHC survival reached ~15% over the most apical 40% of the cochlea.

Conclusion
The small but reliable protective effects of our custom diet are striking given the virtually complete cell death and profound hearing loss observed in control animals. Additional studies are needed to optimize the protective effects observed here, and to further support the long-term goal of translating successful therapeutics from animal models to clinical trials.
Cochlear transfection of X-linked inhibitor of apoptosis protein via round window reduces cisplatin induced hearing loss in guinea pigs

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Background
Previously, we demonstrated that over-expression of X-linked inhibitor of apoptosis protein (XIAP) can delay the development of presbycusis and mitigate noise induced cochlear damage by using transgenic mouse model. Later, we developed and optimized an approach that is minimal invasive for cochlear gene transfection. In this approach, transgene is carried by adeno-associated virus (AAV). The AAV vector is delivered to the cochlea across round window membrane (RWM) which is made permeable to AAV by using a digestive enzyme. In combination with the use of the new generation of AAV with this RWM approach, we fulfill a satisfactory gene transfection in most of the key cell types including outer hair cells.

Methods
In the present study, we examine if the XIAP overexpression induced by cochlear gene transfection via RWM approach can effectively reduce the cochlear damage of a commonly used chemotherapeutic drug, cisplatin, which is applied 10 days after the transfection surgery and at the dose of 4 mg/kg in 3-6 consecutive days. The hearing threshold was tested in auditory brainstem responses (ABR) in closed field and compared in a self-control design between ears with and without XIAP gene transfection.

Results
Gene-transfection via RWM does not significantly damage cochlear function. After cisplatin treatment, both ABR threshold elevation and hair cell loss are significantly less in the ear transfected with XIAP gene. The exogenous XIAP is quantified and compared with endogenous XIAP in Western blotting.

Conclusion
XIAP gene transfection mediated with AAV vectors via RWM approach can effectively reduce the damage of cisplatin on various cell types in cochlea. The reduced cell damage is associated with functional preservation. RWM approach may be a method applicable in human subjects in hearing preservation against ototoxicity.
Hydrogen peroxide production can be measured in lateral line hair cells using HyPer [Tg(myosin6b:HyPer)] expressing zebrafish.

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Background
Oxidative stress caused by reactive oxygen species (ROS) has been hypothesized to play a critical role in hair cell death, however its role in zebrafish lateral line hair cell death is not well described. Characterization of the timecourse of ROS production in response to ototoxic agents would offer insight on the role of ROS, and role of antioxidants in preventing cell death.

Methods
HyPer (Evrogen) is a hydrogen peroxide biosensor that has been used in vivo to detect low levels of intracellular hydrogen peroxide. We have generated a transgenic zebrafish line expressing cytoplasmic HyPer under the control of the hair cell-specific myosin6b promoter [Tg(myosin6b:HyPer)]. Transgenic zebrafish were then exposed to neomycin followed by in vivo imaging at 1 minute intervals using an inverted Marianas spinning disk system. Mitochondrial depolarization was simultaneously examined using tetramethylrhodamine ester (TMRE) labeling. HyPer signal was examined with and without cotreatment with the antioxidant glutathione.

Results
Hydrogen peroxide generation as measured by HyPer fluorescence increased rapidly in response to neomycin. This effect could be blocked by high (1000 µM) but not low concentrations (250-500 µM) of the antioxidant glutathione. Time courses of hydrogen peroxide generation and TMRE signal were measured in response to neomycin. Only high dose glutathione could protect against neomycin-induced hair cell death.

Conclusion
The Tg(myosin6b:HyPer) expressing zebrafish line can be effectively used to study hydrogen peroxide generation in neomycin-treated zebrafish lateral line hair cells. Neomycin-induced hydrogen peroxide generation and neomycin-induced hair cell death could only be blocked by high doses of the antioxidant glutathione in the zebrafish lateral line.

Small molecule inhibitors of cisplatin-induced hair cell death: Results of a 10,000 compound screen in the zebrafish lateral line.

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Background
Cisplatin is a commonly used anti-cancer drug which causes damage to inner ear hair cells and hearing loss. Because cisplatin treatment is given in planned doses, a compound that protects against cisplatin-induced hair cell damage would
be useful for prophylaxis during cisplatin chemotherapy. We sought to identify such a compound using a zebrafish lateral line screening protocol we have successfully used in the past to identify compounds that protect against hair cell death.

**Methods**

We screened a library of 10,000 drug-like small molecules for protection against cisplatin-induced hair cell damage in the zebrafish lateral line. For compounds associated with reduced hair cell damage, dose-response relationships were determined in the zebrafish lateral line using immunocytochemistry. Hair cell counts were performed to quantify the number of hair cells remaining after cisplatin treatment with and without the potential protectant. To evaluate whether the protective compounds altered the chemotherapeutic efficacy of cisplatin, we tested the compounds with cisplatin in cancer cell culture using human lung adenocarcinoma cells from the A549 and NCI-H23 cell lines.

**Results**

Two compounds, CHCP1 and CHCP2, demonstrated dose-dependent protection against cisplatin-induced hair cell death. At the most optimal concentration, CHCP1 and CHCP2 increased hair cell survival (% control) after cisplatin treatment from 37.1 ± 4.0% to 65.4 ± 11.6%, and from 29.7 ± 5.3% to 70.0 ± 10.2%, respectively. The dose-dependent tumoricidal activity of cisplatin was retained in the presence of CHCP1 and CHCP2 when evaluated in tumor cell culture. We also evaluated 5 analogs of CHCP2, and found similar protection among compounds with minor structural modifications.

**Conclusion**

The finding of two small molecules that protect against cisplatin-induced hair cell death affirms the feasibility of screening using the zebrafish lateral line and provides 2 compounds that may have therapeutic benefit as protectants against cisplatin-induced hair cell damage. Further characterization of these compounds and structurally related analogs are being conducted.

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**Copper-induced hair cell death in the zebrafish lateral line requires functional mechanotransduction.**

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**Background**

Copper is known to induce rapid dose-dependent toxicity in zebrafish lateral line hair cells. The mechanism by which copper gains access to the hair cells is not known. Previous studies have suggested a possible role for the mechano-electrical transduction (MET) channel in copper toxicity. Olivari et al. (2008) found decreased copper toxicity in immature hair cells that do not yet have functional MET channels, and decreased toxicity in the presence of the MET inhibitoramiloride. We have examined the effects of chemical and genetic inhibition of mechanotransduction on copper damage in hair cells of the zebrafish lateral line.

**Methods**

Zebrafish larvae were exposed to a treatment condition then incubated in either normal fish embryo media or media with 1 micromolar CuSO4. Tip link proteins, required for normal MET channel function, were disrupted by calcium chelation in calcium-free media with 5 mM EGTA. Quinine was used as a potent chemical inhibitor of the MET channel. Genetic disruption of MET function was assessed using the cadherin-23 mutant, sputnik. After treatment, zebrafish were fixed for immunohistochemistry and hair cell survival was quantified.

**Results**

Consistent with previous studies, we found that copper rapidly induced dose-dependent hair cell death. Tip link disruption with EGTA increased hair cell survival (% control) from 37.8 ± 5.9% to 99.5 ± 7.5%. The sputnik mutant demonstrated hair cell survival of 93.5 ± 8.3%. Quinine partially prevented hair cell death, increasing hair cell survival from 58.9 ± 8.6% to 79.2 ± 6.3%.

**Conclusion**

Chemical and genetic inhibition of mechanotransduction protected against copper-induced hair cell death. This suggests that functional mechanotransduction is important for copper-induced hair cell death in the zebrafish lateral line.
Background
In the therapeutic arsenal in the fight against cancer, chemotherapy has a place. Cisplatin (cis-diamine-dichloroplatinum II [N2Cl2PtH6] or CDDP) is a widely used anti-cancer drug. Unfortunately, it presents a significant toxicity to both the kidney (nephrotoxicity) and the cochlea or inner ear (ototoxicity). Nephrotoxicity of CDDP is circumvented by strategies of hydration, permitting use of increased doses of CDDP. In contrast, no protocol to protect the cochlea is currently available.

The mechanism of action underlying the cytotoxicity of CDDP is based on DNA alterations. In the cochlea, CDDP has been shown to induce an oxidative stress leading to the activation of the mitochondrial apoptotic pathway of the auditory sensory cells. However, very few is known about the upstream actors of the apoptotic events. Those actors could be therapeutic targets to prevent hearing loss in CDDP treated patients.

Methods
To investigate the signalling events between DNA damage and apoptotic death, we use an in vitro model of cochlear primary culture. This model mimics the mechanisms of degeneration observed in vivo after a CDDP treatment, i.e. a dose-dependant toxicity, a greater sensitivity of outer hair cells to CDDP ad a base to apex gradient of degeneration. We used 129/sv mice lacking p53 to understand the role of this transcription factor in vivo when treated with CDDP.

Results
We have shown that, in vitro, p53 expression is upregulated and that p53 is phosphorylated on its serine 15 after CDDP treatment. We also have demonstrated that p53 have a crucial role in the CDDP mediated hearing loss using p53 KO mice. Indeed, the absence of p53 protects from a severe ABR threshold elevation following a cumulative dose of CDDP of twice 8mg/kg.

Conclusion
These results show for the first time that, in vivo, p53 is potential therapeutic target to prevent CDDP-induced hearing loss.
A Combination of Cilostazol and Ginkgo Biloba Extract Protects Against Cisplatin-Induced Cochleo-Vestibular Dysfunction by Inhibiting the Mitochondrial Apoptotic and ERK Pathways

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Background
Cisplatin (CDDP) is an anticancer drug that induces significant hearing loss and balance dysfunction as side effects. Cilostazol (CS) has neuroprotective and antioxidant effects, whereas Ginkgo biloba extract (GbE) has preventive effects on CDDP-induced hearing loss in rats, and GbE enhances the antiatherogenic effect of CS by inhibiting generation of reactive oxygen species (ROS). The purpose of this study was to investigate the effects of renexin (RXN), which contains GbE and CS, against CDDP-induced cochleo-vestibular dysfunction in rats and to elucidate the mechanism underlying the protective effects of RXN on auditory cells.

Methods
The first study was to identify the optimal Renexin dose against cisplatin-induced ototoxicity using adult male Sprague-Dawley rats (n=21). In the second phase, 57 rats were divided into six groups; cisplatin-only, Renexin+cisplatin, Cilostazol+cisplatin, GbE+cisplatin, vehicle+cisplatin and normal control. Rats were intragastrically treated with Renexin during 7 days for pretreatment and 5 days for posttreatment. Cisplatin in dose of 16mg/kg was treated intraperitoneally just two hours after the 7th time of the otoprotective agents. After the 5-day follow-up, ABR test, tail hanging test, and swimming test were performed for hearing function and vestibular function evaluation, respectively. Cochleas and utricles/saccules were harvested for morphological evaluation (SEM). In vitro study, the effect of Renexin on cisplatin-induced cytotoxicity was analyzed in the HEI-OC1 cells. The cell viability, apoptosis, reactive oxygen species generation as well as changes in the signal pathway related to apoptosis were investigated by western blot assay.

Results
Rats intraperitoneally injected with CDDP exhibited an increase in hearing threshold and vestibular dysfunction, which agreed with hair cell damage in the Organ of Corti and otoliths. However, these impairments were significantly prevented in a dose-dependent manner by pre- and co-treatment with RXN, and these preventive effects in RXN-treated rats were more prominent than those in GbE-treated rats. In a CDDP pharmacokinetic study, platinum concentration was very similar between CDDP-only treated and RXN+CDDP co-treated rats. RXN markedly attenuated CDDP-induced intracellular ROS and significantly reduced CDDP-activated expression of p-extracellular regulated kinase (ERK), BAX, cytochrome c, cleaved caspase-3, and cleaved poly (ADP-ribose) polymerase but increased BCL-XL expression.

Conclusion
These results show that RXN may have a synergistic effect by strongly protecting hearing and vestibular dysfunction induced by CDDP by inhibiting ROS production, mitochondrial pathways, and the ERK pathway, without interfering with CDDP pharmacokinetics. Therefore, RXN could potentially be used to reduce CDDP-related hearing loss and dizziness.
The role of connexin 43 and 26 hemichannels in cisplatin-induced apoptosis

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Background
Connexins are assembled into a hemichannel and two identical hemichannels form a gap junction, which play an important role in K+ recycling, metabolic communication in the cochlea. Non-junctional hemichannels can gate open and pass molecules (<1 kDa) by depolarization, extracellular Ca2+, oxidative stress, and metabolic inhibition. The purpose of this study was to investigate the role of hemichannel, connexin 43 (Cx43) and 26 (Cx26) in cisplatin-induced apoptotic process using siRNA techniques.

Methods
We constructed recombinant plasmid of pcDNA 3.1 (-)Cx43, Cx26 and Cx-pcDNA 3.1 (-) plasmid was transfected in Cx-deficient HeLa cells. The formation of functional hemichannels was tested by the transfer of Lucifer yellow (LY, m.w 457.25) from a Ca2+ free condition with/without hemichannel blockers (25μM Carbenoxolone; CBX, 25μM 18 alpha-glycyrrhetinic acid; 18-AGA). To investigate the mechanism for cisplatin-induced regulation of hemichannels, we analyzed opening of hemichannels under 4 different conditions (Ca2+ condition, Ca2+ Free condition, Hydrogen peroxide (H2O2), potassium chloride (KCl) and cisplatin-induced stress condition).

Results
Cx43 and Cx26 are significantly expressed by transfection of pcDNA 3.1(-)-Cx43, Cx26 in the HeLa cells. LY uptake was observed in Cx43, Cx26-transfected cells, but was absent in Cx-deficient wild type HeLa cells and these uptake was inhibited by both CBX, 18-AGA. Under cisplatin toxicity, hemichannels opened in Cx43, Cx26-transfected cells. Cx-transfected cells significantly responded to Ca2+ free condition but little in H2O2 or KCl. Live/dead cell assay and western blot assay indicate that cisplatin-induced cell death in Cx43, Cx26-HeLa cells increased (89%, 66%) compared to wild type HeLa cells. CBX, 18-AGA prevented increase of cell death.

Conclusion
The hemichannel Cx 43 and 26 are opened by cisplatin resulting in apoptosis, which may be closely connected with the suppression of Ca2+. 
Protective Effect of Metformin against Cisplatin Induced Ototoxicity in Auditory Cell Line

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Background
Metformin, an antidiabetic drug with a potent anticancer property, is known to prevent oxidative stress-induced cell death in several cell types through a mechanism dependent on the mitochondria. In the present study, we investigated the influence of metformin on cisplatin ototoxicity in auditory cell line.

Methods
Cell viability was determined by MTT, and oxidative stress and apoptosis were assessed by flow cytometry analysis, Hoeschst 33258 staining, ROS measurement, and western blotting. Intracellular calcium concentration changes were detected with calcium imaging.

Results
Pretreatment with 100uM of metformin prior to application of 20 uM of cisplatin significantly decreased the late apoptosis in HEI-OC1 cells and also, significantly attenuated the cisplatin-induced increase in reactive oxygen species (ROS). Metformin also inhibited the activation of caspase-3 and the expression of poly-ADP-ribose polymerase. Also, pretreatment with metformin prevented the elevation of intracellular calcium concentration induced by cisplatin. We propose that metformin has protective effects against cisplatin induced ototoxicity by inhibiting the increase of intracellular calcium and preventing apoptosis and ROS production.

Conclusion
This is the first study on auditory hair cells to investigate the protective effects of metformin against cisplatin induced ototoxicity. By using HEI-OC1 cells in calcium imaging, we found that metformin inhibited the increase of intracellular calcium, enhanced the cell viability and prevented ROS production.
NAD attenuates mefloquine-induced cochlear damage from reactive oxygen species

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Background
Mefloquine, an anti-malarial drug, is widely used for prophylactic or chemotherapeutic treatment of drug resistant malaria. However, there is growing concern among government agencies about its neurotoxic and ototoxic side effects. Our previous studies demonstrated that mefloquine causes a dose-dependent apoptosis to cochlear hair cells, spiral ganglion neurons and auditory nerve fibers. Nicotinamide adenine dinucleotide (NAD) is a key mediator of important biological processes including energy metabolism, mitochondrial function, calcium homeostasis, and anti-oxidation. Our recent in vitro studies have shown that NAD can protect cochlear hair cells and afferent axons against mefloquine ototoxicity. However, the mechanisms underlying NAD otoprotection are not fully understood. Previous reports indicate that mefloquine increases the production of free radicals leading to cellular damage and apoptosis. These results suggest that the antioxidant properties of NAD may prevent mefloquine-induced cochlear damage.

Methods
We tested this hypothesis by treating cochlear organotypic cultures with mefloquine, NAD or mefloquine plus NAD and measured the resulting cochlear damage and the expression of the superoxide radical in hair cells and neurons using dihydroethidium fluorescence.

Results
When cochlear cultures were treated for 24 h with 35 µM mefloquine nearly all of the hair cells and auditory nerve fibers in the basal turn were destroyed; however, most hair cells and nerve fibers in apical turn were intact. In contrast, when cochlear cultures were treated for 24 h with 35 mM mefloquine plus 20 mM NAD the auditory nerve fibers throughout the cochlear remained intact. When cochlear cultures were treated for 24 h with 50 µM mefloquine, all the hair cells and nerve fibers were completely destroyed in both basal and apical turns. However, many of the hair cells and nerve fibers in the basal and apical turn survived if the cultures were treated with 50 µM mefloquine plus 20 mM NAD. Superoxide expression was assessed 12 h after mefloquine treatment when most cells were still present. Superoxide was heavily expressed in mefloquine treated cochlear explants whereas superoxide was low in cultures treated with NAD alone or mefloquine plus NAD.

Conclusion
NAD suppresses the production of the superoxide radical in mefloquine treated cochlear cultures and provides significant protection against mefloquine ototoxicity.
Lateralization of traveling wave energy in the hearing organs of katydids
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Background
Mathematical models of the curved mammalian cochlea predict that traveling waves propagate along the cochlea by reflection off the lateral walls. Beyond a certain number of reflections, sound waves propagate close to the outer wall analogous to sound waves in whispering galleries due to the cochlea’s graded curvature. This results in lateralization of traveling wave energy. Experimental evidence for this has been lacking as in-vivo measurements of traveling wave propagation in the mammalian cochlea are generally restricted to the basal part along one dimension (longitudinal) due to its poor accessibility. In a previous study it was discovered that traveling waves form the physical basis of frequency discrimination in the hearing organ of katydids, the crista acustica (CA) and that their longitudinal (proximal-distal) characteristics are comparable to those measured in mammals.

Methods
Here we studied in detail the radial (anterior-posterior) structure of traveling waves along the CA in-vivo using laser Doppler vibrometry for stimulus frequencies between 9 and 30 kHz.

Results
During their propagation, sound-induced traveling waves along the CA were reflected off the lateral walls. A clear lateralization of traveling wave energy towards the anterior CA was found. Lateralization of energy had an inverse correlation with radius of curvature. Further, a dynamic rotation of the hearing organ about the longitudinal (proximal-distal) axis was seen during propagation.

Conclusion
Presumably the dynamic rotation and the consequent tilt in the magnitude response generate the input relevant for signal transduction in the auditory sensillum of the CA. Our results are consistent with the predictions made by mathematical models of the curved cochlea. This opens up exciting possibilities of using the CA as a model system to put to test theories of cochlear mechanics.

Two-tone suppression based on somatic motility of the outer hair cell with a cochlear model
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Background
Half a century ago, it was find out that the auditory nerve (AN) firing was suppressed by the other tone. This phenomenon is called two-tone suppression (2TS) and observed in both the basilar membrane (BM) and inner hair cell (IHC). Our interest is mechanisms of the 2TS. However, the mechanisms of the 2TS have been unclear because of technical difficulties. Models can predict the mechanisms of the 2TS. A transmission-line model of the cochlea is much for this problem because of possible to describe internal mechanical and electronically dynamics in the cochlea.

Methods
A transmission-line model of the cochlea containing an outer hair cell (OHC) model was developed in accord with findings about the somatic motility of the OHC. This model was used to investigate how the somatic motility generates the 2TS in the BM.

Results
The 2TS produced by the model was comparable with experimental results in the BM and the AN: (1) The maximum magnitude of suppression exceeded 40 dB around the characteristic frequency (CF) equaled to suppressor frequencies, (2) The distribution of maximum rates of suppression (ROS) was highly correlated with compression in the BM, (3) Maximum ROSs were approximately 2.0 dB/dB for low-frequency suppressors and (4) Maximum ROSs were reduced with increasing suppressor frequency near the CF. The third finding conflicts with BM data. However, the model indicated no filtering effect of the IHC. These results further indicate that the mechanisms of 2TS are based on the somatic motility of the OHC.

Conclusion
The basic mechanism of the 2TS appears to be a reduction in the output of the OHC in response to a probe when the variation in length of the OHC changes from being linear to nonlinear, through an increase in the suppressor level.
Reducing boundary reflection in computational models of the cochlea

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Background

Both the efficiency with which cochlear model responses can be computed and the ease of their subsequent interpretation can often be substantially improved if unwanted reflections from the apical end of the cochlea can be reduced or eliminated.

Methods

We describe and compare several different schemes for reducing boundary reflections in box models of the cochlea. In one promising scheme, based on the concept of perfectly matched layers, the box representing the cochlea is supplemented with a thin boundary layer with modified governing equations or parameters and terminated with a boundary condition. The boundary layer is designed to be absorptive so that the solution decays substantially within the layer and no substantial reflections occur due to the boundary condition. The modified boundary layer is chosen to balance the conflicting needs of achieving a high spatial decay rate while avoiding reflections due to the rapid change.

Results

The modified boundary layer framework is applicable to most traveling wave models, and we illustrate it using passive and active 1D transmission-line models of the cochlea solved in the frequency and time domains. We compare the results with those of other methods, including specifying absorptive boundary conditions, such as terminating with the characteristic impedance.

Conclusion

The modified boundary layer method proves effective at reducing boundary reflections in computational cochlear models. The method is particularly appealing because of its simplicity and its extendibility to other cochlear models and simulation schemes.
Two effects of viscosity revealed by tectorial membrane traveling waves
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Background
The tectorial membrane (TM) is thought to play a critical role in stimulating hair cells as part of a resonant system (Allen, 1980). A key prediction of such models is that subtectorial space damping plays an important role in determining sharpness of tuning. Recently, however, it has been shown that TM traveling waves (Ghaffari et al., 2007) may contribute to cochlear tuning by longitudinally coupling radial cross sections. Here we test the relationship between viscosity and tuning for TM waves and compare that relation to results for wave and resonance models.

Methods
We measured wave properties of mouse TMs immersed in artificial endolymph solutions with poly-ethylene glycol (PEG) added to increase viscosity. Two PEGs with different molecular weights (MW) were used: one (8 kDa) chosen to penetrate TM pores (Masaki et al., 2006) while the other (400 kDa) could not.

Results
Our findings show that introducing small MW PEG increases TM wave speeds by \(\sim 40\%\) and decreases wave decay constants by \(\sim 35\%\). Analysis of a lumped parameter model of the TM showed that these changes in wave parameters can be explained by a change in shear viscosity from 0.2 Pa*s to 0.65 Pa*s with no accompanying change in shear modulus. In contrast, introducing large MW PEG has little effect on wave speed (\(-2\%) or decay (\(-9\%), suggesting shear viscosity inside the TM is significantly more important compared to fluid viscosity.

Conclusion
Our results suggest that fluid surrounding the TM has two separate effects on TM motion. First, increasing fluid viscosity has the obvious effect of increasing drag on the surface of the TM. In addition, if high viscosity fluid penetrates the TM’s porous structure then the TM’s material properties are also directly affected. Of these two effects, we see that the latter has a much greater impact on TM waves.

Interestingly, these results counter a fundamental assumption in classical cochlear models: that increasing damping would broaden tuning. By contrast, scaling symmetry suggests that the decrease in wave decay constants caused by increased viscosity would sharpen tuning. These results highlight important differences between wave and resonance models of the TM and offer entirely new ways to think about cochlear tuning.
Acoustic Emissions from Coupled Strings
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Background
Linear mode conversion is a phenomenon that has been studied for several decades and finds application in plasma physics, geophysics, and auditory mechanics. Typically, one propagating wave can excite another at a location where the wavelengths of the different modes become similar and energy can be exchanged. The slowly varying WKB approximation becomes singular at such a mode conversion point. Here we examine a simple mechanical system that illustrates that effect. Of particular interest is the understanding of how a reflection can occur when there are no discontinuities in physical properties. Originally, Rayleigh considered this problem on a single string with smoothly varying mass over a finite transition region. The reflected waves are an analog for stimulated emissions from the ear.

Methods
We consider traveling transverse waves on two identical uniform taut strings that are elastically coupled through springs that gradually decrease their stiffness over a region of finite length. The wave system can be decomposed into two modes: an in-phase mode (+) that is transparent to the coupling springs, and an out-of-phase mode (-) that engages the coupling springs. The system exhibits linear mode conversion whereby an incoming (+) wave is reflected back from the smooth transition both as a propagating (+) wave and an evanescent (-) wave, while both types emerge as propagating forward through the transition. We match a local transition layer expansion to the WKB expansion to obtain estimates of the reflection and transmission coefficients.

Results
The oscillatory behavior of the reflection coefficients with respect to the frequency parameter shows that even this simple system exhibits a stimulated emissions spectrum that is characteristic of the ear. In the present system the frequency spacing of the spectrum originates from the oscillatory nature of the Airy functions. Physically, it is due to the coupling stiffness gradient. In the mammalian ear there are numerous contributions to a decreasing coupling stiffness gradient in the organ of Corti. The increasing length of outer hair cells and their stereocilia from base to apex are an example. In contrast, the coherent reflection theory that Zweig and Shera developed for the ear argues that the incoming wave is scattered by local irregularities of any kind, and then coherently filtered by the incoming wave.

Conclusion
This work was supported by grant DC000033-17 from the Intramural Program of the National Institute of Deafness and Other Communication Disorders.
High Sensitive Inertia Sensor Mimicking Stereocilia Array
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Background
Stereocilia could achieve high sensitivity over a broad dynamic range by the interplay of adaptation and negative stiffness mechanism, which is the limitation of conventional cantilever-type inertia sensors due to the mass and volume constraint. To overcome the conventional sensor design limitation, in this study, we proposed a high sensitive inertia sensor mimicking hair bundle's amplifying mechanism.

Methods
The proposed sensor consists of 10 sensing units of inverted pendulum array, pairs of magnet, which is controlled by stepping motor along the guide bar that mimics the readjustment of tip link tension by sliding molecular motors. The translation of magnets along the guide bar moves inverted pendulum to lean side way in a gradual manner. When the restoring force of the spring at the pivotal joint exceeds the maximum magnetic repulsive force, an abrupt change of the direction of pendulum lean occurs. Ten magnets on the guide-bar have slight phase differences, depending on the position of the magnet on the guide-bar and a moving direction, which would then induce differences in sensitivity among pendulum arrays. The count of abrupt change of pendulum position is measured by photo detector. To examine the characteristics of the proposed sensor, we measured the force-displacement relation and sensitivity when step and sinusoidal input is applied to the system.

Results
The inverted pendulum unit shows negative stiffness near the origin due to the magnetic repulsive force. The negative stiffness region is shifted when stepping motor moves the magnet on the guide-bar side-to-side. We observed the spontaneous oscillation which was induced by the interplay between the negative stiffness and adaptation mechanism. Experiment results showed that the count of abrupt change of pendulum position was proportional to applied force, implying the proposed biomimetic system could serve as an inertia sensor. Nonlinear compressive sensitivity was observed at the small force input compared to the conventional sensor.

Conclusion
We proposed a biomimetic inertia sensor which overcomes the limited sensitivity issue of conventional engineering sensors. The current system is limited in its range of measurable force due to constraints of the peak oscillation magnitude, which should be further resolved by biomimetic sequential resetting mechanism.
Simulation of tuberculoventral inhibition in the cochlear nucleus and its role in enhancing tone responses in the cochlea through the olivocochlear reflex pathway

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Background
Wickesberg and Oertel (1988) discovered that tuberculoventral (TUB) cells in the dorsal cochlear nucleus (DCN) provide inhibitory post-synaptic potentials to bushy and multipolar cells in the ventral cochlear nucleus (VCN). The inhibition was found to be delayed and frequency-specific (Wickesberg and Oertel, 1990) and may help achieve echo cancellation.

Methods
In the present work, a computer model is built to simulate the TUB inhibition of T-multipolar (TM) cells. The model allows both TUB and TM cells to be phase-locked to tonal stimuli at sufficiently low frequency. The model has also been integrated with models of the cochlear mechanics (Liu and Neely, 2010), the inner hair cells and the auditory nerves (Meddis 1986; Sumner et al., 2002), and the MOC efferents (Liu et al., 2012), so dynamics of the entire network can be calculated.

Results
Simulation shows that, if the delayed inhibition reaches its maximum strength at time 1/CF (characteristic frequency), the inhibition can reduce tone responses in the TM cells but not their noise responses.

Conclusion
Because the TM cells project to medial olivocochlear (MOC) interneurons, the TUB inhibition in the cochlear nucleus may work to enhance the cochlea’s response to tones in noise by reducing the MOC reflex strength in a spatially confined manner.

Stereocilia lipid membrane: Nonlinear hair-bundle mechanics and channel activation

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Background
The Gating Spring Response, calculated from hair bundle force-displacement measurements, behaves in a non-linear manner. The measurement shows a region of increased compliance that overlaps with mechanoelectric transduction (MET) channel gating. The presented computational work investigates the potential role of the lipid bilayer at the ciliary tip complex in providing the physical basis of: nonlinear mechanics in bundle activation; MET channel opening; and the correlation between them.

Methods
The first part of the biophysical model describes motion of the bundle by rigid body kinematics implementing pivot motion of a cilium with respect to its base and shear motion between adjacent cilia through the sliding mechanism of horizontal top connectors. The second key part of the model describes the elastic deformation of the lipid bilayer at the tip complex. The continuous lipid membrane of the cilia is partitioned into two regions with functionally distinct properties. The model
describes the elastic flexure and stretching of the cap region under the action of the tip link force while accounting for lipid molecule diffusive transport between the cap and body regions. In this model, the body region acts as a lipid reservoir for the cap region and differential molecular diffusivities of these two regions are postulated to be crucial to the membrane tensioning response. Finally, using the lipid bilayer free energy density near the tip-link lower insertion, a Boltzmann function was used to calculate an open probability of the MET channel. With all components coupled, the model is capable of simulating the potential impact of the lipid at the tip complex for the MET channel opening.

Results
There are two key results from the model simulation. First, nonlinear bundle force-displacement can be generated by the lipid membrane tension without correlating contribution from the MET channel. The consistency of the simulation results to the experimental measurements suggests the existence of an isolated tip lipid compartment with a radial size of about 25nm, likely defined by membrane-skeleton-crosslinkers attachment. Second, MET channel can be activated by the free energy generated by the curvature on the tensioned membrane.

Conclusion
The proposed modeling framework demonstrates the plausibility of the lipid bilayer serving as the underlying mechanism responsible for the gating spring. It also indicates that the force transfer through the lipid can provide the gating energy needed to open the MET channel.

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Estimation of Added Fluid Mass to Vibrating Cochlear Partition and its Contribution to the Frequency-Place Relation of Mammalian Cochlea
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Background
A pure tone vibration delivered to the oval window travels along the cochlear partition (CP) until it culminates at a location specific to the tone. The traveling wave has been explained with mechanical models represented by a series of spring-mass resonators interacting with the fluid of cochlear ducts. The frequency-dependent peak response locations are primarily determined by the stiffness and mass of the CP. The stiffness gradient of most cochlear models is about 11 dB/octave, which is greater than experimentally measured stiffness gradients (4-9 dB/octave: Naidu & Mountain, 1998; Emadi, Richter et al., 2004). We hypothesized that the added fluid mass of the CP due to its fluid interaction, increases toward the apex so that a broader frequency range can be encoded with a limited stiffness gradient.

Methods
A finite element (FE) model of the CP based on anatomical and mechanical measurement data of gerbil cochlea was used to estimate the mass of the CP. Its static responses were validated against experimental results (Nam & Fettiplace, 2010). From the dynamic response of the FE model without fluid-interaction, we obtained the effective mass of the CP that depends on the geometry and the vibrating mode. In order to evaluate the fluid mass, a Neely's 1D passive cochlear model (1981) was adopted with stiffness and mass of spring-mass resonators obtained from the FE analyses. Due to added fluid mass, the peak resonating frequency at a location was lower than the natural frequency of the CP spring-mass resonator. The fluid mass was obtained from the difference between those two frequencies.

Results
The stiffness of the CP ranged from 760 to 0.3 mN/m per 10 µm section for 30 to 0.16 kHz frequency range (9 dB/oct). The mass of the CP per 10 µm section ranged from 30 (base) to 60 ng (apex), while the fluid mass ranged from 5 (base) to 1000 ng (apex). The structural mass was dominant only at the most basal locations (< 2mm from the basal end).

Conclusion
Except at the most basal locations, the effect of added fluid mass is greater than the structural mass of CP which agrees with a previous study (Lim & Steele, 2000) and recent experimental measurement (Dong & Olson, 2009). The stiffness gradient dominates the frequency gradient near the base while fluid mass gradient explains the frequency gradient near the apex.

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Simulations of two tone suppression and distortion products in a physiologically-based computational model of the cochlea

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Background
Two tone suppression and distortion products are well known characteristics of the response of the cochlea to two tones. We have recently developed a nonlinear physiologically-based computational model of the cochlea (Biophys J., 2012, 102:1237-1246) and demonstrated that the model reproduces most aspects of the response of the cochlea to a single tone (frequency tuning, compressive nonlinearity, harmonic distortion and DC shift). We extend the previously developed model to the simulation of two tone interactions in cochlear mechanics.

Methods
In the model nonlinear hair bundle mechanoelectrical transduction is coupled to outer hair cell somatic electromotility. The structural model includes degrees of freedom for the basilar membrane (BM) and tectorial membrane (TM) with longitudinal coupling and is coupled to a three-dimensional representation of the fluid. The nonlinear model is formulated in the frequency domain and is solved using an iterative alternative frequency-time scheme. The response of the cochlea to two tones is decomposed into carefully chosen harmonic components. Using forward and inverse Fourier transforms, the nonlinearity of the transduction current is introduced in the time-domain while the equations are solved in the frequency domain.

Results
Model simulations of two suppression and distortion products on the BM are compared to experimental data. Simulations reproduce the effect of a suppressor tone on the magnitude and phase of the BM response. The spatial extent of suppression on the BM response is analyzed. For distortion products simulations, the spectrum of the response of the BM to two tones is presented. The dependence of the cubic distortion product on the level and frequencies of the primaries is compared to measurements.

Conclusion
The physiologically-based computational model reproduces many aspects of the nonlinear response of the cochlea to two tones. These simulations support the theory that cochlear amplification is mainly due to somatic electromotility and that cochlear nonlinearity arises from the nonlinearity of hair bundle mechanoelectrical transduction.

Carhart's notch: A window into compressional and inertial mechanisms of bone-conducted hearing

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Background
It is well known that otosclerosis is accompanied by a sensitivity decrease in the 1-2 kHz range for bone-conducted (BC) hearing. This now clinically referred to as Carhart's notch because of its importance in the clinical assessment of this pathological condition. However, the mechanism for Carhart's notch remains to be poorly understood. We developed a computational framework to address this question.

Methods
A three-dimensional finite element (FE) model of a human middle ear coupled to a cochlea was formulated. The geometry of the middle ear and cochlea, including semicircular canals, was obtained from micro-computed tomography (µCT) images. In this study, BC hearing was simulated by: (1) applying rigid-body vibrations for the inertial mechanism; and (2) pressure on the outer bony shell to investigate the effects bone compression mechanism. The stapes annular ligament was stiffened to simulate otosclerosis in the model. The change in hearing by the BC route was calculated by the difference in the maximum basilar membrane (BM) velocity between the normal and the otosclerotic conditions.

Results
Applying only inertia to the FE model caused about 20 dB hearing loss at low frequencies (below 0.5 kHz) which is not consistent with the clinical measurements. The inertia caused 30 dB hearing loss at 1.3 kHz, which decreased to 0 dB as frequency increased to 10 kHz. On the other hand, applying only bone compression caused no hearing loss at all calculated frequencies (0.2 kHz - 10 kHz) such that there was no notch in the 1-2 kHz frequency range. Applying both inertia and bone compression stimulation removed the hearing loss at low frequencies (0.2 - 0.3 kHz) returning to normal
BC hearing. Applying both conditions also showed a 25 - 35 dB hearing loss in the 1 - 2 frequency range which was qualitatively consistent with clinical measurements.

Conclusion
The compressional mechanism of BC is dominant at 0.5 kHz and below and is unaffected by fixation of the footplate to the bony wall. At frequencies above 2 kHz, the inertial component of BC is dominant. Between 0.5 to 2 kHz, the dominant factor can be either inertia or bone compression depending on the magnitude ratio between two components. The magnitude ratio can differ between individuals due to the different outer bony shell thickness, degree of stapes fixation, or mass of middle-ear ossicles (or fluid in the cochlea).

Directional Inertia at the Base of the Cochlea and Bone-conduction Hearing

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Background
For the most part, analytical and numerical models have not demonstrated any functional importance for the coil shape of the cochlea for air-conducted (AC) hearing. We hypothesize that this coiled shape affects the bone-conducted (BC) route of hearing by inertial forces of the middle ear and cochlear fluid which can be tested by stimulation from different directions.

Methods
A three-dimensional finite element (FE) model of a human middle ear coupled to the inner ear was formulated. The geometry of the middle ear and cochlea, including semicircular canals, was obtained from micro-computed tomography (\(\mu\)CT) images. BC excitations were simulated by applying rigid-body vibrations normal (in the orthogonal direction) to the BM at 0.8 (\(d_1\)), 5.8 (\(d_2\)), 15.6 (\(d_3\)) and 33.1 (\(d_4\)) mm from the stapes such that relative motions of the fluid within the cochlea produces excitations of the basilar membrane (BM). A second component of the input comes from the relative motion from the mass inertia of the ossicles.

Results
The vibrational direction normal to the BM surface at the base \(d_1\) of the cochlea produced the highest BM velocity response across all tested frequencies - higher than an excitation direction normal to the BM surface at the non-basal locations (\(d_2\)-\(d_4\)) and their best-frequency (BF). The basal part of the human cochlea features a well-developed hook region, in which the BM undergoes a sudden curvature that produces the largest difference in fluid volume between the scala vestibuli (SV) and scala tympani (ST) not found elsewhere in the cochlea. Due to this sudden curvature, the normal direction to the BM surface in the base differs significantly from the normal directions to the BM along the mid-turns and apical regions of the cochlea. This change caused the most significant anti-symmetric pressure difference than the other direction so that the BM velocity is maximized for the \(d_1\) location rather than vibrations at the other locations.

Conclusion
The proximity of the hook region to the oval and round windows, combined with it having the biggest fluid volume and impedance difference between the SV and ST, results in maximization of the pressure difference between the SV and ST for BC stimulation perpendicular to the BM in this region, and consequently the resulting BM velocity to be maximized.

A Finite Element “Virtual Labyrinth” Model of the Human Inner Ear Facilitates Electrode Design for a Multichannel Vestibular Prosthesis

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Background
Accurate modeling of prosthetic electrical current flow within the human labyrinth can facilitate optimal design of electrode arrays for a multichannel vestibular prosthesis intended to restore sensation to individuals with bilateral loss of vestibular hair cell function. Our previous research has demonstrated the ability of a finite element and neuromorphic computational model to accurately predict responses to electrical stimulation delivered by vestibular implant electrodes in chinchillas and
rhesus monkeys. We extended this approach to human anatomy to create a virtual labyrinth model of the human vestibular system.

Methods
Model geometry was developed in Amira® from the Fudan University Virtual Temporal Bone Data, a 3-dimensional (3D) reconstruction of a human specimen serially sectioned at 50 um. Electrodes were virtually positioned within the model volume to test hypotheses regarding dependence of neural responses on electrode location, stimulus current and insulating materials displacing inner ear fluids. Finite element analysis (ComsolMultiphysics®) computed the extracellular potential field time course during current pulses from any constellation of electrodes. Extracellular potential waveforms then served as inputs to stochastic, nonlinear dynamic models for 2,415 vestibular afferent axons with spiking dynamics based on a modified Smith and Goldberg model incorporating parameters that varied with fiber location in each endorgan. Action potential propagation was implemented by a well validated model of myelinated fibers. 3D eye rotation axes were predicted from the relative proportion of model axons excited within each of the three ampullary nerves.

Results
Consistent with model predictions in rodents and nonhuman primates (in which measured 3D vestibulo-ocular reflex [VOR] responses were well-predicted by model output), the magnitude and direction of predicted 3D VOR responses depend heavily on electrode proximity to cristae (changing significantly for displacements on the order of 100 um), return electrode location, and the presence of insulating material (such as fat or silicone) displacing perilymph and/or endolymph. Current spread and the degree of spurious activation of nontarget afferents was similar to that observed in simulations and empiric data for rhesus monkeys.

Conclusion
Although model validation must await empiric 3D VOR and electrophysiologic measurements in humans implanted with multichannel vestibular prostheses, the success of analogous models for rodents and monkeys suggests the Virtual Human Labyrinth can facilitate optimal design of electrode arrays intended for human implantation.

In vitro culture of adult mouse cochlea
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Background
Study of mammalian inner ear for hair cell regeneration, biophysics, genetic deafness and drug screening has been greatly hampered by the lack of a system that enables the culture of adult cochlea in vitro. In standard culture condition, adult mammalian cochleae rapidly lose all hair cells, with the degeneration of overall structure of sensory epithelium, leading to downregulation of supporting cell gene expression, further loss of supporting cells, and overgrowth of fibroblasts.

Methods
Here we developed a protocol with which the integrity of cultured adult mouse cochlea can be maintained in vitro. With the protocol, a relatively intact adult cochlear sensory epithelium structure could be preserved for 10-14 days in vitro.

Results
Supporting cells in culture were well organized with a distinct cellular pattern similar to that in vivo. Supporting cells were labeled with multiple specific markers, supporting that the identity of supporting cells was maintained. Many inner hair cells, especially those in the apex, survived while outer hair cells were lost during culture. In contrast to traditional attachment culture system, there was a minimum contamination from other cell types including fibroblasts within the sensory epithelium. When infected by adenovirus carrying a GFP reporter, the infected supporting cells and inner hair cells started to express GFP, which could be observed two days after infection, with the expression maintained for 1 to 2 weeks. While outer hair cells were rapidly lost within 12 hours in normal culture, the death of outer hair cells can be substantially delayed by blocking necrosis pathway.

Conclusion
Our in vitro culture system thus provides a new tool to investigate the adult mammalian cochlea through genetic manipulation and pharmacological interference.
Hypotonic Condition Induces Hearing Impairment In TRPV4 Knock-out mice

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Background
Changes of osmolality in the inner ear fluids have been considered to cause some clinical disorders such as tinnitus, fluctuating hearing loss, and Mendier's disease. Transient receptor potential vanilloid 4 (TRPV4) has been recently proposed as an osmo- and mechanosensitive channel and reported to be involved in cellular volume regulation and affect cellular function. In the cochlea, TRPV4 was confirmed to express in the outer hair cells, inner hair cells as well as spiral ganglion cells. To investigate the influence of osmotic pressure on cochlear function, we investigated whether the hypotonic condition in the inner ear fluids causes hearing impairment in TRPV4 knock-out mice.

Methods
Hypotonic solution was introduced into the inner ear of mouse through the left side of posterior semicircular canal by an osmotic mini-pump (Alzet®: 1003D). Wild type (C57BL/6) and TRPV4 knock-out (KO) mice were implanted with the pump and cannula for delivery of artificial perilymph (300 ± 2mOsm) or hypotonic saline (150 ± 2mOsm) continuously for 3 days. Auditory brainstem responses (ABRs) were measured on day0, day3 and day7 after the surgery at the pump-implanted side. Both sides of cochleae were removed at day7, and then processed for further histological assessment.

Results
At day7, mice implanted pump with artificial perilymph showed less than 10dB threshold shifts at all frequencies. Wild type mice showed statistically significant smaller ABR threshold shifts compared to KO mice at all frequencies 7 days after the implantation of pump with hypotonic solution. The non-pump-implanted (contralateral) side showed significant higher OHC survival rate than the pump with hypotonic solution implanted (experimental) side in KO mice at day7. On the other hand, difference of the IHC survival rate was not significant.

Conclusion
Hypotonic condition induced hearing impairment in TRPV4 knock-out mice which was statistically significant compared to that of the wild type mice. It indicates that TRPV4 might be involved in hearing impairment by changing of osmolality in the inner ear fluids. Any disturbance in the homeostasis of inner ear fluids may thus affects the functional properties of OHCs via TRPV4, thereby causing some hearing impairment.
**Auditory System in Knock-in Mouse Models of Pendred Syndrome**

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**Background**

Among all persons with congenital hearing loss, 4–10% are cases of Pendred syndrome (Morton and Nance, 2006), which is an autosomal recessive disorder. This syndrome is caused by mutations in the pendrin gene. Mutations in pendrin also cause nonsyndromic hearing loss (Usami et al., 1999). Pendrin is a membrane protein and expressed in the inner ear. In the inner ear, pendrin is involved in the conditioning of the ion concentration of the endolymph. Generally, membrane proteins are transported to the plasma membrane; however, some protein mutants are retained in the cytoplasm. In the case of pendrin mutants, the amount of pendrin that localizes on the plasma membrane is low, and the ion concentration of the inner ear is abnormal. So far, no effective therapy for hearing loss caused by these mutations has been established. In this study, to clarify the mechanisms of the Pendred syndrome or nonsyndromic hearing loss caused by the H723R pendrin mutant, which is a major cause of sensorineural hearing loss among Japanese, and to find a curative method for hearing loss in humans, a knock-in mouse model with H723R pendrin mutant was developed as a first step.

**Methods**

Two strains of knock-in mice, namely, mice carrying the knock-in point mutation of H723R pendrin (murine-type knock-in mice) and mice in which the mouse pendrin was replaced by the human H723R pendrin mutation (human-type knock-in mice) were generated using a recombinneering approach. The hearing ability of these mice was assessed using the ABR test.

**Results**

In this study, the hearing of thirty-three murine-type knock-in mice was assessed. Twenty-one of them had hearing thresholds of less than 40 dB SPL (Fig. 1) and twelve showed mild to moderately severe hearing loss (hearing thresholds of 40–70 dB SPL) at various ages of onset (Fig. 2). All human-type knock-in mice at the age of 4 weeks had a profound hearing loss (> 90 dB SPL).

**Conclusion**

It is suggested that the murine-type knock-in mouse model may be comparable to cases of human patients since various shift patterns of hearing among patients with mutations in the pendred gene have been reported (Napiontek et al., 2004). There is a possibility that hearing loss of the mice with early onset was caused by H723R mutation in the pendred gene. Human-type knock-in mice had profound hearing loss. This result is comparable to studies on knock-out mouse models (Wangemann et al., 2007).
Optimizing a $2f_1$-$f_2$ DPOAE Measurement for Extended High Frequencies.

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**Background**

Given the variation in cochlear mechanics from base to apex, it is not surprising that different stimulus conditions are necessary to generate the most robust distortion at different frequencies. Specifically, sharper tuning (due to reduced spread of excitation) would dictate that narrower frequency ratios between the stimulus tones ($f_1$ and $f_2$, $f_2 > f_1$) used to elicit $2f_1$-$f_2$ DPOAEs would be necessary to produce optimal overlap between mechanical excitation patterns. Typically, the stimulus parameters that elicit the most robust DPOAEs up to ~8 kHz in normal-hearing ears are used in clinical applications (Gorga et al. 1997). Commonly used (standard) parameters include $f_2/f_1=1.22$ and stimulus levels of 65/55 dB SPL. However, Dreisbach and Siegel (2001) have shown the optimal frequency ratio ($f_2/f_1=1.22$ at 1 kHz) decreases with increasing frequency (~1.16 at 10 kHz). Additionally, DPOAEs are most sensitive to hearing loss when recorded using low or moderate stimulus levels; however, DPOAEs are more difficult to obtain at higher frequencies using these levels. Taken together, this body of evidence suggests that stimulus parameters used routinely at frequencies <8 kHz are suboptimal for measurements at higher frequencies. This study aimed to optimize DPOAE measurements up to 20 kHz using stimulus parameters chosen to account for underlying variations in the mechanical properties over the length of the cochlea to further enhance our ability to detect cochlear damage.

**Methods**

Accurate and stable measurements of DPOAE amplitudes using custom hardware and calibration procedures for swept-(up to 16 kHz; n=15) and discrete-frequency (up to 20 kHz in ~1/8-octave steps; n=15) stimuli were obtained in normal-hearing individuals. An innovative test paradigm, comprising three fixed-level combinations and varying $f_2/f_1$ ratios in comparison to a “standard” protocol and/or a “fixed” ratio protocol was evaluated.

**Results**

Larger DPOAEs were measured with narrower frequency ratios than the standard ($f_2/f_1=1.22$) at frequencies ≥6 kHz in normal-hearing ears. Increasing stimulus level elicited larger DPOAEs (most notably at higher frequencies). Evaluation in 15 ears with hearing loss revealed no compromise to the sensitivity of DPOAEs as a screener for mild hearing loss.

**Conclusion**

Preliminary results support the hypothesis that using stimulus conditions optimized for variations in presumed underlying cochlear mechanics are needed to elicit robust DPOAEs at higher frequencies. Further evaluation is needed to explore whether DPOAE test performance improves when optimal stimuli are utilized.
Investigating stimulus-frequency otoacoustic emissions and threshold fine structure at high frequencies
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Background
When the inner ear is stimulated by a tone, energy at the stimulus frequency propagates back out to the ear canal via mechanisms that depend on active cochlear processes. These stimulus-frequency otoacoustic emissions (SFOAEs) can be extracted from the ear canal signal using techniques that exploit cochlear nonlinearity. Additional evidence for backwards-propagating energy within the cochlea can be found in the quasiperiodic fluctuations in hearing sensitivity, termed threshold fine structure. This fine structure pattern is thought to arise from standing waves produced by multiple reflections of energy between the middle-ear boundary and the site of SFOAE generation. The high frequency limits of the mechanisms underlying SFOAE generation and threshold fine structure have not been thoroughly explored; here we report both acoustic and behavioral evidence for these mechanisms extending beyond 8 kHz.

Methods
Behavioral hearing thresholds and SFOAEs were obtained from adult participants in an exploratory fashion. Pure-tone detection thresholds for frequencies between .125 and 20 kHz were measured using a modified Bekesy tracking procedure, with frequency steps ranging from 1 to 1/100th of an octave. SFOAEs were evoked by both swept and discrete tones for probe frequencies between .4 and 18 kHz, and extracted from the ear canal pressure via either the compression or suppression method. Stimuli were calibrated using a method that compensated for the depth of probe insertion in the ear canal. Spontaneous OAEs were identified via spectral averaging of the ear canal signal obtained without stimulation.

Results
Threshold fine structure was observed at frequencies that approached the start of the steeply sloping high-frequency portion of the audiogram, which typically ranged from 11-14 kHz, even in an ear with a broad mid-frequency region of elevated thresholds. The depth of the high-frequency fine structure was often comparable to that found at lower frequencies (~10 dB). Robust high-frequency SFOAEs were also measurable in some ears when elicited by discrete tones, with amplitudes peaking around 10-13 kHz, and reliable phase gradients measurable up to ~15 kHz. Swept-tone SFOAE measurements above 10-12 kHz were complicated by excessive variability in the pressure at the probe frequency, most likely due to small changes in probe position across stimulus repetitions.

Conclusion
The mechanisms underlying SFOAE generation and threshold fine structure extend to the highest frequency regions of sensitive hearing. Further work is needed to determine precisely how SFOAE amplitudes are related to fluctuations in threshold.
Latency of Tone-Burst-Evoked Auditory Brainstem Responses and Otoacoustic Emissions: Level, Frequency, and Rise-Time Effects

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Background
Auditory brainstem responses (ABRs) and otoacoustic emissions (OAEs) are two physiological measures that have been used to indirectly estimate basilar membrane (BM) delays. Simultaneous measurement of ABRs and OAEs may provide insights into effects of level, frequency and stimulus rise time on BM delay.

Methods
Tone-burst-evoked ABRs and OAEs (TBOAE) were measured simultaneously in 46 normal-hearing human subjects. Total data-collection time was 620 hours. Stimuli included a wide range of frequencies (0.5 to 8 kHz, ½-octave steps) and levels (20 to 90 dB SPL, 10-dB steps). Multiple tone-burst rise times at each frequency allowed characterization of the dependence of latency on rise time. ABR latencies were estimated by three judges and averaged. TBOAE latencies were estimated as the group delay of the OAE waveform. In an effort to facilitate comparisons between ABR and TBOAE latencies, the level-dependent component of ABR latency (i.e., forward latency) was derived by subtracting level-independent estimates of synaptic and neural delays (5 ms) from wave-V latency.

Results
ABR latencies showed the same orderly frequency and level dependence as previously reported [Gorga et al., J. Speech Hear. Res. 31, 87-97 (1988)]. The level-dependence of TBOAE latencies was similar to that of ABR latencies across all frequencies, although the TBOAE latencies were more variable by a factor of five. Additionally, the frequency dependence of TBOAE latencies was similar to ABR latencies only for mid-frequencies. Rise time also had a significant effect on ABR latency, but this effect was independent of its level dependence. Comparison between simultaneously-measured ABR and TBOAE latencies showed that TBOAE latency is about twice ABR forward latency for frequencies above 1.5 kHz, but the two latency estimates were roughly equal for lower frequencies.

Conclusion
The level dependence of ABR latency must be either mechanical or neural, whereas TBOAE latency is determined by cochlear mechanics. Thus, the similarity at all frequencies of the level dependence of ABR latency to that of OAE latency supports the view that most of this level dependence is due to cochlear mechanics. This view is further supported (1) by independence of level and rise-time effects and (2) by similarity of frequency dependence at mid-frequencies. [Work supported by the NIH R01 DC2251 and P30 DC4662].
Towards a clinical test for the medial-olivocochlear reflex (MOCR)

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Background
A clinical test for the medial-olivocochlear reflex (MOCR) must be fast, reliable, valid, and the MOCR strength measured by the test must vary widely in the population. Such a test must also be as free as possible from confounds, such as multiple OAE components, middle-ear muscle reflexes (MEMR), negative middle-ear pressure, and spontaneous OAEs.

Methods
In a series of experiments, we have developed and evaluated (and continue to refine) three OAE-based MOCR tests based on transient-evoked OAEs (TEOAEs), stimulus-frequency OAEs (SFOAEs), and most recently swept-frequency OAEs (SWOAEs). All three measure the reflection-component OAE that is more sensitive to the MOCR.

Results
Our SFOAE and TEOAE-based tests take around 3 minutes per ear to run, with repeated testing needed to assure stability. Avoiding the MEMR is nigh-on impossible in many ears - it is both difficult to reliably measure in many ears and its magnitude is often at the same level as the raw MOCR effect. But for SFOAE and SWOAE tests we can show which measurements are affected by this confound.

Conclusion
All methods have their pros and cons, with SWOAEs showing the most potential, but needing more development.
Binaural Measurements of Otoacoustic Emissions in Humans
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Background
The efferent system has been shown to be involved in the protection of the cochlea against acoustic trauma and the facilitation of signal detection from background noise, but the precise function is still controversial. Due to the fact that the efferent fibers innervate the outer hair cells (OHCs), otoacoustic emission (OAE) measurements are a suitable tool to assess efferent function. Especially distortion-product OAEs (DPOAEs), which are evoked by two-tone stimuli are often used to study cochlear mechanics. Two types of DPOAEs are distinguishable, the cubic distortion tone (2f1-f2) and the quadratic distortion tone (f2-f1). The present project investigates efferent effects on cubic and quadratic DPOAEs under binaural stimulation using the same two-tone stimuli in both ears, which should produce a natural hearing impression.

Methods
Thirteen subjects participated in two measurement series. OAEs were recorded using two identical Etymotic ER-10C probe systems. Part I: Level-growth-functions in the level range of 10 to 70 dB SPL were measured at 6, 4, 2 and 1 kHz during binaural and monaural two-tone stimulation to evoke cubic DPOAEs. The f2/f1 ratio was adjusted to 1.2. The measurements were repeated with levels up to 80 dB SPL at 6 and 4 kHz with the individual optimal ratio to evoke quadratic DPOAEs. Part II: The recordings on the cubic and quadratic emissions were repeated with repetitive amplitude-modulated two-tone complexes.

Results
Three types of efferent effects occurred in the subjects. (i) Level-dependent suppression in one ear versus enhancement of DPOAE levels in the other ear, resulting in a strong lateralization. (ii) Level-dependent suppression or enhancement in both ears and (iii) effects at DPOAE level maxima or minima. In some cases, only one ear showed suppression and the other ear was not affected by binaural stimulation. The type of effect could change when instead of pure tone stimuli amplitude-modulated pure tones were used. The effects were independent of the stimulus frequency, but highly level-dependent, with largest changes of up to 4-6 dB in the low-level stimulus ranges. The effects on the quadratic emissions were less stable than the ones on the cubic emissions, but had the same magnitude.

Conclusion
The results indicate a different processing of pure tone and amplitude-modulated sound signals already at the level of the cochlea, which may be mediated by a shift in the OHC’s operating point.

Estimates of Human Cochlear Tuning Derived from DPOAE Reflection-Component Delays
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Background
Past work has shown a consistent relationship between reflection emission phase-gradient delay and cochlear tuning in a variety of mammalian species, as predicted by filter theory and models of otoacoustic emission (OAE) generation (Shera et al., 2010). Here, we exploit this relationship and explore methods of estimating delay trends, with the long-term aim of studying cochlear tuning throughout the human lifespan.

Methods
The 2f1-f2 DPOAE was recorded with pure tones (L1,L2 = 65,55 dB SPL, f2/f1 = 1.22) swept at 8 s/oct in 186 subjects from prematurely born neonates to elderly adults. DPOAEs were measured from 0.5–4 kHz in all groups and extended to 8 kHz in young adults. DPOAE reflection components were separated via inverse FFT and their phase-gradient delays calculated and expressed in periods. Following parametric study to optimize variables, trend lines were fit to group data using two strategies: energy-weighted loess and a loess fit to segments at amplitude peaks only. A mean trend combining both approaches was also computed. Tuning ratios derived from measurements in non-human mammals and shown to be species invariant were applied to the human data to generate measures of filter bandwidth.

Results
The phase-gradient delay versus frequency function shows a typical bend (putative apical-basal transition) between 0.8 and 1.2 kHz, consistent with past OAE work. Qerb estimates from young adults roughly match those obtained from SFOAEs and ranged between 10–15 for frequencies below 4 kHz, increasing to approximately 17–18 for higher
frequencies. The infant groups tend to show sharper tuning than the adult subjects over much of the frequency range. The middle-aged and elderly groups manifest greater variability in Qerb across frequency, including a slight decrease at the highest frequency tested.

**Conclusion**

DPOAE reflection components can provide reliable estimates of delay and tuning. The match with SFOAE-based tuning estimates is imperfect for predictable reasons: a) The strength of the probe generating reflection at 2f1-f2 is not controlled in the DPOAE paradigm and undoubtedly varies among individuals, increasing the variance of the tuning estimates; b) F1 in the DPOAE paradigm produces suppression at the DP place (Kalluri and Shera, 2001), influencing reflection generation. Despite these caveats, the Qerb values derived here approximate those reported by Shera and colleagues (2010) in humans and represent a first step towards quantifying the maturation and aging of human cochlear tuning.

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### Aging of the Medial Olivocochlear Reflex: Associations with Speech Perception in Noise

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**Background**

The medial olivocochlear reflex (MOC) has modulatory effects on cochlear mechanics, typically producing inhibition by hyperpolarizing outer hair cells. Otoacoustic emissions (OAEs) can gauge this reflex. Little is known about how MOC effects change throughout the human lifespan. Here, we studied aging effects on the MOC reflex using an expanded repertoire of indices while considering the effects of MOC activation on both the 2f1-f2 distortion product OAE and on its dual components. Also, we preliminarily examine the links between age-related changes in medial efferent effects and speech perception in noise.

**Methods**

DPOAEs were recorded with pure tones (L1, L2 = 65, 55 dB SPL; f2/f1 = 1.22) swept at 8 s/oct in 116 subjects, with and without contralateral acoustic stimulation (CAS) to activate the MOC reflex. Age groups included teens and young, middle-aged and elderly adults. The MOC reflex was calculated as the vector difference between baseline and +CAS conditions at fine structure peaks only, to minimize the effects of component interference. Additionally, the DPOAE was separated into distortion and reflection components via inverse FFT and the effects of CAS on the amplitude and phase of each were calculated. The middle ear muscle reflex (MEMR) was monitored during testing. Speech perception was assessed under headphones using a variety of speech tokens, for a range of signal-to-noise ratios.

**Results**

CAS presented at 60 dB SPL produced a reduced contralateral MOC reflex in middle-aged subjects compared to teens and young adults. In general, MOC activation reduced emission magnitude most strongly at low frequencies and tended to produce shallower reflection-component phase slope. Unexpectedly, the elderly group showed the largest reductions in emission level, and over half of these subjects also showed a pronounced increase in DPOAE fine structure when contralateral noise was presented, most notably at 65 and 70 dB SPL. Links between these findings and speech perception in noise are currently being probed.

**Conclusion**

By middle age, the medial efferent reflex is weakened; however, the oldest subjects show seemingly incongruous effects of MOC activation. Both observations in the elderly -stronger MOC reflexes and increases in DPOAE fine structure- might be explained by unintended activation of the MEMR during presentation of the contralateral noise elicitor. Middle ear muscle contractions could increase multiple internal reflections, producing the pronounced fine structure noted here; they might also increase the size of the MOC reflex by reducing DPOAE levels during reverse transmission.

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### The Half-Octave Shift Revisited: Evidence from Middle-Ear Power Transmittance and Stimulus Frequency Otoacoustic Emissions in Humans

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**Background**

Following exposure to an intense tone, the largest temporary otoacoustic emission level shift (TES) occurs approximately one-half octave above the exposure frequency. Variation in TES between individuals may be partially explained by
individual differences in middle-ear sound transmission. The aim of this study was to examine the contribution of middle-ear sound transmission to measured amounts of TES.

**Methods**

Pre-exposure measurements of middle-ear power transmittance (MEPT) and stimulus frequency otoacoustic emissions (SFOAE) at 1.4 and 2.0 kHz were obtained from normal-hearing adults. SFOAE levels were derived from probe alone and probe plus suppressor conditions. SFOAE measurements were repeated as a check of test/retest reliability. Following a 1-minute exposure to a 105 dB SPL tone at 1.4 kHz, SFOAE measurements at 1.4 and 2.0 kHz were completed at 2 and 4 minutes after the exposure. Shifts in SFOAE levels at 2 minutes (TES$_2$) and 4 minutes (TES$_4$) were calculated.

**Results**

The mean SFOAE test/retest reliability was within 1 dB at both 1.4 and 2.0 kHz. The mean amount of TES was greater than one standard deviation above the mean value of test/retest reliability only at 2.0 kHz. The amount of TES$_2$ and TES$_4$ was similar at both test frequencies. Trends in the data indicated that MEPT was related to the amount of TES. Individuals with higher MEPT values at 1.4 and 2.0 kHz tended to have larger TES$_4$ at 2.0 kHz, while individuals with lower MEPT values tended to have smaller TES$_4$ at 2.0 kHz.

**Conclusion**

The results of this study are consistent with previous research indicating the maximum effect of tonal overexposure occurs one-half octave above the exposure frequency. For SFOAEs, the amount of TES$_4$ appears to be related to a) MEPT at the exposure frequency and b) MEPT at one-half octave above the exposure frequency. These findings suggest that MEPT of intense sound may play a role in explaining some of the variability seen in the amount of TES across individuals. Exploration of the role of MEPT in temporary threshold shift is warranted.

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Otoacoustic-Emission Detection when the Middle Ear Contains Amniotic Fluid: A Chinchilla Animal Model

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**Background**

Otoacoustic emissions (OAE) have frequently been used for newborn hearing screening. However, OAE screening tests have a low specificity and have referral rates higher than recommended by the Joint Committee on Infant Hearing. The presence of residual amniotic fluid in the middle ear is one of the reasons for these problems. The aim of this study was to determine the effects of human amniotic fluid on distortion product OAEs and noise levels across various frequencies. The effects of normal saline were also studied in order to investigate the role of viscosity.

**Methods**

Forty-six adult female chinchillas with normal Preyer's reflex were used. The animals were initially screened using auditory brainstem responses and tympanometry. Only animals with normal hearing and middle-ear status were included. They were randomly divided into 8 groups based on the type (amniotic fluid or saline) and volume (0.5, 1, 1.5, 2 ml) of liquid introduced into the middle ear. OAE measurements were taken at the Montréal Children’s Hospital using a SmartOAE system (Intelligent Hearing Systems) under inhalational anaesthesia. After an initial OAE measurement, liquid was introduced into the middle ear by trans-bullar injection and then a second OAE measurement was made.

**Results**

Both the type and the volume of liquid made a difference in both the OAE amplitudes and the noise level. Significant reductions of OAE amplitudes occurred across all frequencies, with more reduction for greater volumes. Changes in the noise level had important effects on the signal-to-noise ratio at some frequencies.

**Conclusion**

Both human amniotic fluid and saline in the middle ear of a chinchilla animal model resulted in changes in OAE detection patterns that may be relevant to newborn hearing screening with OAEs.
Exploring stimulus conditions for measurement of the quadratic f2-f1 distortion product otoacoustic emission
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Background
Measurement of distortion product otoacoustic emissions (DPOAEs) aids in diagnosis of sensorineural hearing loss. Clinically, the cubic DPOAE 2f1-f2 is most commonly employed. However, quadratic DPOAEs (e.g., f2-f1) may also be sensitive to outer hair cell physiology. The clinical utility of DPOAEs may increase if f2-f1 can be used to pinpoint subtle cochleopathies, including those that result from striavascularis damage. Previous work by Bian and Chen (2008) suggests optimal stimulus parameters for f2-f1 differ from those used to elicit robust 2f1-f2 DPOAEs for an f2 of 4 kHz. Here, we explore the optimal f2/f1 ratio for eliciting f2-f1 over a wide frequency range in normal hearing and hearing impaired adults.

Methods
Fifteen young adults (18-30 years) with clinically normal hearing (0.25-8 kHz) in at least one ear participated in this study. In addition, a small cohort of subjects with hearing loss were also examined. The DPOAE at f2-f1 was tested at six f2/f1 ratios (1.14, 1.18, 1.22, 1.30, 1.32, and 1.36) for f2 frequencies ranging from ~660-19,220 Hz. Sweep tones (L1,L2=70 dB SPL) were used to elicit emissions. In addition, behavioral hearing thresholds were obtained from 125-20,000 Hz in the test ear. DPOAE data were averaged in 1/3-octave bins to assess the effect of f2/f1 ratio on f2-f1 amplitude across frequency. The influence of high frequency hearing thresholds on f2-f1 amplitude was also explored.

Results
In contrast to previous reports (Bian and Chen, 2008), these data revealed that f2/f1 ratio was not highly influential in determining f2-f1 level. Although wider ratios did occasionally produce more robust emissions in individual subjects, mean data do not support a ratio or frequency-dependent trend. In some subjects, prominent 2f1-f2 DPOAEs could be measured while little, if any, f2-f1 could be recorded. Conversely, f2-f1 was more robust than 2f1-f2 in some individuals. Hearing impaired subjects demonstrated reduced f2-f1 compared to the normal hearing group.

Conclusion
The f2/f1 ratio did not have a significant impact on mean f2-f1 emission amplitude, with the exception of some individual differences. These preliminary results suggest that the DPOAE at f2-f1 may provide information about cochlear health that is complementary to that obtained from the 2f1-f2 DPOAE.
Estimating Cochlear Frequency Selectivity with Stimulus-frequency Otoacoustic Emissions
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Background
Each point along the basilar membrane of the cochlea is tuned to a specific frequency. At low stimulation levels this mechanical filtering is sharpened due to action of the outer hair cells (OHC). It has been suggested that the tuning of the cochlear filters can be derived from measures of otoacoustic emissions – byproducts of OHC action. Two approaches have been proposed for estimating cochlear frequency selectivity using single frequency OAEs (stimulus-frequency OAEs), based on (a) SFOAE group delays (GDs) and (b) SFOAE suppression tuning curves (STCs). We aim to compare these estimates of frequency selectivity with more direct measurements of cochlear tuning – compound action potential tuning curves (CAP TCs) – obtained in the same animal.

Methods
Twelve chinchillas served as experimental subjects. The CAP and SFOAE STC data were obtained at up to four probe frequencies (1, 4, 8, and 12 kHz). SFOAE STCs were measured as iso-response curves where the level of a suppressor tone is adjusted until a predefined change in ear canal pressure at the probe frequency is recorded. An analogous paradigm was used for CAP TC measurements. The SFOAE GDs were measured using the suppression method for probe frequencies ranging from 0.3 to 12 kHz. The SFOAE data were collected at probe levels of 30 – 40 dB SPL and CAP data for tone-burst levels resulting in a clear response above the noise floor (25 – 60 dB SPL). Q10 values were calculated for all tuning measures.

Results
On average there was good agreement between Q10 derived from SFOAE STCs and CAP TCs at all frequencies except 1 kHz, where emission tuning curves were considerably broader. Q10 derived from SFOAE GD were in good agreement with CAP data at 1 and 12 kHz while for mid frequencies SFOAE GD indicated sharper tuning. When compared to published Q10s for auditory nerve (AN) TCs, CAP TC and SFOAE STC seemed to underestimate sharpness of tuning, while the opposite trend was observed for SFOAE GD Q10s. When individual data were analyzed there was no clear one-to-one positive relationship between Q10s derived with any pair of the three methods.

Conclusion
Although, on average, the sharpness of tuning derived with all methods changed in a similar way across the frequency range, the individual data indicate that neither SFOAE measure allows for precise estimation of CAP tuning within a given subject.

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The influence of compression on evaluating dynamic changes in frequency selectivity in forward masking
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Background
Frequency selectivity has been reported to increase as a function of the duration of a forward masker [e.g. Bacon and Jesteadt, J. Acoust. Soc. Am. 82, 1926-1932 (1987)]. This finding suggests that the tuning of the auditory system may sharpen shortly after being stimulated; however, this interpretation does not consider the influence of nonlinearity in the cochlea. The current study tests the hypothesis that an increase in frequency selectivity with masker duration is a product of cochlear nonlinearity. This was done by measuring frequency selectivity psychophysically and correcting these measurements with an estimate of cochlear nonlinearity.

Methods
Frequency selectivity was estimated from “input filter patterns” (IFPs) where the detection threshold for a 4-kHz probe was measured in the presence of a masker whose level was held constant and whose frequency varied. Cochlear nonlinearity was estimated from on- and off-frequency growth of masking (GOM) and “output filter patterns” (OFPs) were obtained by transforming IFPs by this estimate.

Results
Preliminary results suggest that corrected measurements of frequency selectivity (i.e. OFPs) do not differ between long and short maskers. This suggests that compression may be responsible for the increased frequency selectivity observed in IFPs measured with a long masker.

Conclusion
These results can be interpreted as suggesting, 1) that the tuning of the auditory system does not change during stimulation, or 2) that manipulating masker duration may be an insensitive way to study how auditory tuning changes with stimulation. The influence of the medial olivocochlear reflex on frequency selectivity supports the latter interpretation as this reflex has been reported to reduce frequency selectivity when elicited [e.g. Cooper and Guinan, J. Physiol. 576, 49-54 (2006)].
Heart Rate Responses of Big Brown Bats (Eptesicus fuscus) to Conspecific Vocalizations

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Background
Acoustic communication plays a primary role in social interactions among many species of bats. Our previous work revealed a complex acoustic communication system among big brown bats in which acoustic cues and call structure signal the emotional state of the vocalizing animal. We found that heart rate of a calling bat scaled to the intensity level of evoked vocal aggression, confirming our behavioral state classifications assessed by vocalizations and behavioral displays. However, it is not known how these social vocalizations modify the behavior and affective state of a listener.

Methods
In this study we monitored the heart rate responses of big brown bats (n=4) to the presentation of conspecific vocalizations. An external heart rate monitor was attached and signals were transmitted to a wireless receiver (TBSI). Bats were allowed to roost freely within a small cage, in which heart rate, video, and vocalizations were recorded over 5-min trials. Bats were presented with sound stimuli that corresponded to one of five different contexts: high aggression, lower aggression, appeasement, and neutral pure tones at two different intensity levels. Stimuli were presented 30 times at a rate of 1/s, and the sound context of trials was randomized within each recording session.

Results
Bats generally showed a significant increase in heart rate in response to sound (81%, 57 trials), including pure tones. However, we found that the conspecific vocalizations evoked a greater increase in heart rate than pure tones (p<0.05). Surprisingly, lower aggression vocalizations evoked the largest change in heart rate magnitude (p<0.001) and remained elevated for longer durations (p<0.001) compared to any other sound context. Heart rate responses also showed level dependence, with the higher-intensity tones evoking a larger increase in heart rate than lower-intensity tones (p<0.005).

Conclusion
The results of the current study suggest that factors influencing heart rate changes in response to conspecific vocalizations may be different for the sender and receiver. Lower aggression vocalizations, which evoked the greatest increase in heart rate of receivers, are more likely to be associated with close physical contact and biting. In contrast, vocalizations of a highly aggressive bat are more likely to prevent physical contact and reduce the probability of biting. The increased probability of biting associated with lower aggression vocalizations may explain why receivers show a greater elevation in heart rate in response to these vocalizations. Supported by NIDCD grant R01 DC00937.

Two-step Cluster Analysis for Automated Classification of Mouse Pup Isolation Syllables

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Background
Acoustic communication signals can be used as behavioral markers of stress or diseased state. For example, mouse pups vocalize at higher rates when they are cold or isolated from the nest. Beyond rate of calling, recent work also suggests that changes in the types or proportion of emitted syllables can also carry significant information about the internal state of the animal. However, current methods of manual syllable categorization are time consuming and vulnerable to experimenter bias.

Methods
In this study we use an automated two-step clustering technique to determine the number of statistically distinct syllable types produced by CBA/CaJ mouse pups, and compare the results to prior manual classification methods. A total of 15,116 CBA/CaJ mouse pup syllables were used to determine the number of distinct syllables within the repertoire at different ages (p5, 3,145 syllables; p7, 4,329; p11, 4,560; p13, 3,082). For each syllable, an automated 9-point fundamental frequency contour was first computed. Next, the distribution of frequency transitions was analyzed to develop a criterion for indicating discontinuous frequency steps. The two-step cluster analysis was computed using four variables for each syllable: the total number of frequency steps and the start, middle, and end frequency of the fundamental.

Results
Manual classifications of these vocalizations had previously identified ten syllable types based on spectro-temporal features (Grimsley et al., PLoS One, 6 (3), 2011). The automated cluster analysis identified four distinct clusters: two syllable types with continuous frequency-time structure that were distinct based on their frequency bands, and two syllable
types that featured single or multiple discontinuous frequency transitions. Although the cluster analysis computed fewer syllable types than manual classification, the clusters are more acoustically distinct and represent the probability distributions of the acoustic features within syllables.

Conclusion
Using fixed categorical boundaries informed by the cluster analysis, we generated a Microsoft Excel-based mouse syllable classification calculator that rapidly classifies syllables, with over 90% correspondence. This tool can be used by behavioral neuroscientists to investigate the relationship between mouse pup vocal signals and genetic makeup, health status, or emotional state. Supported by NIDCD grant R01 DC 00937.

Measurement of the Frequency Dependence of Medial Olivocochlear Activation During an Auditory Intensity Discrimination Task
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Background
In awake humans in passive listening (i.e. not doing a task), tones and narrow-band noise evoke medial olivocochlear (MOC) efferent activation over broad frequency regions with the largest MOC inhibition not necessarily centered on the frequency of the presented sound. Recently, it has been shown that during an intensity discrimination task, MOC activity is increased compared to passive listening to the same sounds. Since the MOC efferents have been postulated to facilitate stimulus intensity discrimination in noise, the task-related increase in MOC activity might be near the frequency of the target tone where it might best help the auditory discrimination. With this in mind, we sought to determine whether task-related MOC activity is restricted to, or correlated with, the target frequency of the task.

Methods
Human subjects with normal hearing are recruited for a two-interval forced choice intensity discrimination task. In each trial, two 20 ms tone pips at the same frequency, separated by 380 ms, are embedded in a broadband noise and presented to one ear per subject. The task is to indicate whether the two tone pips are of the same intensity, or not. MOC activity is monitored by click-evoked otoacoustic emissions (CEOAEs), in the task ear before and immediately after the task, and, sometimes, in the contralateral ear also during the task. The decrease in CEOAE level, relative to the level at the beginning of the task, is taken as an index of the MOC inhibition. In each subject, two target frequencies (>1 octave apart) are used with sets of trials at one target frequency alternated with sets of trials at the other target frequency. For comparison, subjects are also tested using the same sounds but without doing a task.

Results
Preliminary results indicate that the pattern of MOC activation across frequency shows a dependency on the task frequency.

Conclusion
The paradigm has great potential for showing the extent to which the cochlear frequency region receiving MOC inhibition follows the task frequency. We will have more complete results at the ARO meeting.

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Relationship between middle-ear transmission characteristics and frequency modulation detection

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Background
The frequency-modulation difference limen (FMDL) with a low modulation rate has been used as a measure of the listener's sensitivity to the temporal fine structure (TFS) of a stimulus, which is represented by the pattern of neural phase-locking at the auditory periphery. However, factors that determine inter-listener variation of FMDL have not been thoroughly explored. This study examined the extent to which non-neuronal factors, specifically mechanical middle ear transmission characteristics, can account for inter-listener variation of FMDL. The middle-ear transmission characteristics and the FM detection performance in normal-hearing listeners were compared.

Methods
The middle-ear transmission was assessed by wideband energy reflectance measurement, in which the reflectance and admittance are derived from the sound reflected by the middle ear. The reflectance and admittance frequency response were measured as a function of frequency from 0.2 to 6 kHz. FMDLs were measured by using a 2I-2AFC transformed up-down method. A stimulus was a 750-ms long tone burst with a frequency of 1 kHz. The modulation rate was 2 Hz. Sixteen audiometrically normal adults participated in the experiment as listeners. In the analysis, participants were divided into two groups: the lower-FMDL group (nine listeners) and the higher-FMDL group (seven listeners).

Results
The reflectance was typically near 1.0 (all of the sound energy was reflected) at low frequencies (<0.5 kHz). The reflectance-versus-frequency function exhibited a minimum at frequencies within a range between 1 and 4 kHz (the resonance frequency). The resonance frequency of the higher-FMDL group was distributed around 1 kHz, while that of the lower-FMDL group was distributed above 1 kHz. Consistently, the admittance value at 0.25 kHz (which is generally assumed to be negatively correlated with the stiffness of the middle ear; a decrease in the stiffness is associated with a lower resonance frequency), exhibited a significant correlation with FMDL (Pearson's $r=0.57$, $p=0.02$).

Conclusion
The results showed that an appreciable fraction of inter-listener variability in FMDL could be accounted for by the listener's middle ear characteristics: FM detection performance tended to be poor when the stimulus frequency was close to the listener's resonance frequency. When an FM detection task is used to evaluate the degree of neural phase-locking, the inter-listener variation of middle-ear transmission should be considered.

Auditory Cortex is Highly Tuned to the Emergence of Regular Patterns in Sound Sequences

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Background
We used psychophysics and magnetoencephalography (MEG) functional brain imaging to assess listeners’ ability to detect the emergence and violation of complex regularities (characterized by long repeating patterns) in ongoing sound sequences and the degree to which this process is bottom-up driven or dependent on explicit attention. Stimuli were tone pip sequences that contained transitions between random and regular frequency patterns. Transitions from a regularly alternating to a random tone sequence (REG-RAND) are immediately detectable as the first tone to violate the established regularity pattern is sufficient to signal the transition. In contrast, listeners must wait longer (at least one regularity cycle) to detect the opposite transition—from a random to a regular pattern (RAND-REG).

Methods
We used sequences of 50ms tone pips arranged according to 4 frequency patterns: REG sequences consisted of a regularly repeating pattern of X tones ($X=10, 15, 20$; new pattern for each trial). RAND sequences consisted of a sequence of tones of random frequencies. REG-RAND and RAND-REG sequences contained a transition between a regular and a random pattern. Additionally a proportion of CONTROL stimuli consisting of a simple step change in frequency between two long pure tones were used to estimate basic response times (RT). In all signals, the time of change was jittered across trials.

In the behavioural experiments ($N=16$), subjects were actively detecting the transitions in REG-RAND, RAND-REG and CONTROL stimuli (change occurred in 50% of the trials). The time required to detect the emergence/violation of regularity was estimated by subtracting the RT to CONTROL signals from that to RAND-REG and REG-RAND. In the MEG
experiment (N=16; different subjects), naïve participants listened to RAND-REG and REG-RAND stimuli while performing an unrelated visual decoy task.

Results
Behavioral response times reveal that subjects required about a cycle and a half to detect the emergence of regularity in RAND-REG signals. Since the transition is not detectable before the first regularity cycle, our results show that listeners only required an additional half cycle to recognize the onset of regularity. Analysis of MEG data is ongoing. Preliminary results indicate that brain response latencies (when the subjects were not actively listening to the transitions) reliably match the RT data.

Conclusion
Our data reveal that the auditory system is remarkably efficient at detecting the appearance and disappearance of regularities in sound sequences, even for very long patterns.

A perceptual measure of cochlear gain control by the medial olivocochlear system
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Background
The medial branch of the efferent olivocochlear (MOC) system controls the gain of the cochlear amplifier. This gain-control function is thought to play an important role in speech-in-noise perception. Otoacoustic emissions (OAEs) offer a qualitative measure of the MOC system’s effect on cochlear gain, but a quantitative measure is still lacking. The aim of the current study was to develop such a measure.

Methods
Cochlear gain affects the ear’s input-output (I/O) function, that is, the function relating the size of the ear’s response to the intensity of the input stimulus. The I/O function can be measured behaviourally by measuring the temporal masking curve (TMC) of a given stimulus. The TMC measures the forward-masking effectiveness of the stimulus as a function of time after stimulus offset. Here, we measure the effect of contralateral noise, which is known to activate the MOC system, on the TMC. In order to control for any longer-lasting effects of MOC activation, the conditions with and without contralateral noise were presented either in a blocked or interleaved sequence. For comparison, we also measured the effect of contralateral noise on OAEs under the same conditions.

Results
The TMCs exhibited a significant effect of contralateral noise, suggesting a gain reduction of ~6 dB. This effect was observed equally in the blocked and interleaved sequences. The OAE also showed a significant effect of contralateral noise (of 0.5 dB, on average), but, in this case, the effect was observed only for the interleaved sequence. There was also no correlation across participants between the sizes of the contralateral noise effects on the TMCs and OAEs.

Conclusion
Our results suggest that TMCs are sensitive to cochlear gain control through the MOC system. However, the lack of correspondence with OAE estimates of MOC-mediated gain control suggests that the two methods are sensitive to different aspects of MOC function. The differences between the two methods may be related to the influence of attention; in the TMC measurements, participants were engaged in an active task, whereas the OAEs were measured under passive listening conditions. Under this assumption, TMCs would appear to offer an ecologically more valid measure of MOC function than OAEs.

Specific Deficits of Basic Auditory Processing in High-Functioning Pervasive Developmental Disorders
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Background
Individuals with pervasive developmental disorders (PDD), such as high-functioning autism (HFA) and Asperger's syndrome (AS), often have difficulties perceiving speech in noisy environments. Here, we investigated whether this might be explained by deficits in basic auditory processing.

Methods
We conducted a set of psychophysical experiments assessing various aspects of basic auditory processing for a group of young adults with PDD (N=21; 5 AS, 9 HFA, and 7 PDD-NOS) and a group of age/IQ-matched controls (N = 37). None of them had hearing levels > 20 dB at all audiometric frequencies. The psychophysical measures included: (a) auditory filter shape at 500 and 2000 Hz estimated by the notched-noise method, (b) level increment detection threshold for noise, (c) gap detection threshold for noise, (d) frequency discrimination threshold for tones at 100 and 1100 Hz, (e) detection threshold for the shift of frequencies of otherwise harmonic components; the stimulus containing harmonics with intermediate resolvability with a fixed spectral envelope, (f) discrimination threshold of interaural time difference (ITD) for noise, (g) discrimination threshold of interaural level difference (ILD) for noise, (h) binaural masking level difference, (i) discrimination threshold of Huggins pitch, and (j) speech reception threshold (SRT), defined as the speech-to-noise ratio at which approximately 50% of Japanese monosyllables are correctly identified.

Results
Statistically significant differences in the median and distribution of psychophysical measures between the PDD and control groups were found for the discrimination thresholds of ITD (f) and ILD (g), as well as SRT (j). In these three tasks, the performance for the PDD group was worse than those for the control group. For the frequency-shift detection task (e), where the temporal fine structure of sounds was the major cue, the difference in the median between the groups was not significant but that in the distribution was significant. In this task, the proportion of good and bad performers was also significantly different between the groups. No statistically significant difference was found for other tasks.

Conclusion
The results indicate that a certain proportion of individuals with PDD have reduced ability to detect temporal fine structure and binaural disparities of sounds, which may contribute to the speech-in-noise perception difficulties experienced by them. The deficits may be related to the malformation of the superior olivary complex found recently in autistic spectrum disorders.

Multiple sound sources in an Australian amphibian?
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Background
Terrestrial vertebrates are capable of producing a wide variety of sounds. Among these are vocalizations which are generated by the larynx in the case of amphibians, reptiles and mammals, or by the syrinx in birds. While the syrinx has the ability to produce separate sound components on each side of its bipartite structure, the larynx is not so well endowed, being restricted to generating one sound at a time that is then subjected to post-filtering by the structures inside the head. Nevertheless, the calls of the Cascade Tree Frog (*Litoria pearsoniana*) exhibit clear examples of simultaneous production of multiple vocalization components.

Methods
All field recordings of this species were made between 2000 and 2200 h from 5 to 10 February, 2012. Males of *L. pearsoniana* were found in their natural habitat in Springbrook, Queensland, Australia, calling along streams and near waterfalls. Vocalizations of seven males of *L. pearsoniana* were recorded with a broadband, directional microphone (G.R.A.S. 40 BE) and preamplifier (G.R.A.S. 26 CB) and a portable digital recorder (Sound Devices 722) at 48 kHz sampling rate. Before each recording, it was determined that no other calling males were in the vicinity of the focal male, thus ensuring that the recorded vocalizations were all coming from a single individual. Data were saved as .wav files and analyzed (FFT, 1024), and displayed using SELENA, a custom-designed program (S. Andrzheevski, St. Petersburg).

Results
Sound spectrograms of the advertisement calls of males of *L. pearsoniana* revealed that the advertisement calls consisted of multiple harmonic stacks in which the fundamental frequencies ranged between 0.5-0.6 kHz, and exhibited various degrees of FM. Multiple instances of simultaneous occurrences of two call components were found. These were most
often seen when one harmonic stack disappeared abruptly and a second stack with a new fundamental frequency would commence before the first stack ended.

Conclusion
One interpretation of our findings is that the observed “stack overlaps” reflect the existence of a second sound source in this frog species. The vocal cords within the larynx are one source; the other is unknown. Several potential candidates for the second source are discussed.

Probing the Auditory Attention Filter Along the Auditory Pathway
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Background
Evidence that auditory attention can be focused in a particular frequency range was discovered by Greenberg and Larkin (1968) using the probe-signal method. Listeners were presented with a cue tone and asked to detect a subsequent tone embedded in noise whose frequency was either the same as the cue (signal tone) or different (probe tones). For tones within 150 Hz of the cue-tone frequency, performance was better for the signal tone than for nearby probe tones, suggesting that listeners focused attention on the cue frequency with an “attentional filter”. However, the mechanism by which such filtering arises is unknown. Moreover, for the just-detectable tones used, listeners were unaware of the signal and probe tone frequencies, thereby excluding a purely cortically mediated explanation. The current study uses the probe-signal method along with physiological measures of auditory response (otoacoustic emissions and EEG) to investigate responses at different levels along the auditory pathway.

Methods
Subjects were presented with three intervals: a cue interval, consisting of a clearly audible cue tone embedded in a 450-ms noise, followed by two 450-ms noise bursts, one of which was randomly selected to contain a near-threshold test tone. The test tone was either presented at the same frequency as the cue tone (75% of the trials) or at a frequency within 150 Hz of the cue frequency (25% of the trials). Performance was analyzed as a function of test tone frequency and then compared across trials differing in the cue-tone frequency. Click trains were presented immediately before and after each three-interval trial to simultaneously record click-evoked otoacoustic emissions (CEOAEs). Additionally, electrical potentials (EEG) from the scalp were recorded using a 32-electrode cap, enabling measurement of both brainstem and cortical responses to the stimuli.

Results
Physiological measures were recorded during the attention task so that their amplitudes could be directly related to task performance. The signal-to-noise ratio for the brainstem and cortical correlates were optimized using principal component analysis across the 32 channels. The CEOAE spectra in the pre- and post-trial conditions were used to assess peripheral changes during the task.

Conclusion
Simultaneous recording of CEOAEs and EEG signals during a behavioral task allows probing of related physiological responses at different stages of the auditory pathway. We expect that a full analysis of the data will elucidate peripheral, sub-cortical, and cortical correlates of the auditory attention filter.
Glycinergic Inhibition Controls Synaptic Integration in the Medial Superior Olive

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Background
The brainstem nucleus of the medial superior olive (MSO) encodes differences in the arrival time of sounds between the ears (interaural time difference or ITD), providing information about the location of sounds in the horizontal plane. MSO neurons receive bilateral phase-locked excitation and glycinergic inhibition. MSO neurons integrate all four synaptic inputs, but it is not understood how these inputs interact at the cellular level to generate ITD output functions. Here, we investigated whether the precise timing of inhibition is important for synaptic integration in the MSO.

Methods
We performed patch-clamp recordings of MSO neurons in acute slices from adult (postnatal day 60 – 90) Mongolian gerbils (Meriones unguiculatus) at near physiological temperature (35 °C). We investigated the interaction between EPSPs and IPSPs using fiber stimulation and conductance-clamp recordings. We also created a computational model of an MSO neuron, based largely on our measured membrane properties.

Results
We first measured excitatory and inhibitory synaptic conductances evoked by putative single presynaptic fibers in voltage-clamp in many neurons. From these data, we selected representative conductance waveforms as templates and then simulated excitatory and inhibitory potentials (EPSPs and IPSPs) in conductance-clamp. We found that the peak of a combined EPSP and IPSP is advanced when the EPSP occurs during the hyperpolarizing phase of an IPSP, but is delayed when the EPSP occurs during the re-depolarizing phase of the IPSP. This suggests that precisely timed inhibition can shift the peak timing of excitation arising from either the ipsilateral or contralateral side. We went on to investigate how the timing of an IPSP influences the summation of two EPSPs. Without inhibition, two EPSPs summed maximally when they occurred simultaneously. However, when a pair of EPSPs occurred during the hyperpolarizing phase of the IPSP, maximal summation was observed when the first EPSP led the second. Moreover, when the EPSPs occurred during the re-depolarizing phase of the IPSP, maximal summation was observed when the first EPSP trailed the second. We also used the same conductance templates as inputs for our model simulations and found that if the model quantitatively matches the voltage responses, it provides a good prediction for the effect of inhibition on the peak timing and summation of EPSPs.

Conclusion
Our findings indicate that precisely timed inhibition can influence the coincidence detection of ipsilateral and contralateral excitation. This provides evidence for a crucial role for inhibition in ITD tuning in vivo.
Cross-correlation based Characterization of Axonal Delays in the Nucleus Laminaris of the Barn Owl using a Microelectrode Array

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Background
The barn owl localizes sound sources in the azimuthal plane by measuring the Interaural Time Difference (ITD). The proposed mechanism for the calculation of these ITDs involves axonal delay lines. The properties of these structures, such as axonal conduction velocities, have been a subject of research for many years.

Methods
In this study, we analyzed data collected in vitro using a Microelectrode Array (MEA) from barn owl chicks aged from postnatal day 2 to day 13. The aim of the analysis was to verify the existence of the axonal delay lines and to characterize the conduction velocities. Improvements with respect to previous studies were made in the preprocessing of the electrophysiological data leading to a higher signal-to-noise ratio and better temporal precision. Furthermore, an algorithm for the determination of the conduction velocity based on cross-correlation was developed and employed. The new algorithm allowed the simultaneous and unsupervised measurement of conduction velocities across the MEA involving recordings from multiple electrodes.

Results
Using these methods, the data was found to show a structured delay distribution consistent with the existence of delay lines. The conduction velocities obtained were lower than reported from in-vivo experiments in mature barn owls as well as from similar in-vitro experiments in chicken.

Conclusion
One factor causing the lower conduction velocities may be that the delay lines are still developing at this age. The methods used in this study might be applicable to future MEA experiments and experiments using multiple simultaneous recordings in general.

Glycine Receptors Expressed in "Timing" Neurons of the Avian Auditory Brainstem Modulate GABAergic Inhibition

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Background
Glycine (GLY) transmission is prominent in the mammalian sound localization circuit, but has not been a focus of investigation in the avian brainstem for several reasons. First, inhibitory input is completely blocked by application of the GABA_A receptor antagonist bicuculline in the nuclei known to process temporal features, nucleus magnocellularis (NM) and nucleus laminaris (NL). Additionally, GLY immunoreactivity appears much less robust compared to GABA staining in these nuclei. However, recent studies from our group and others have shown that GLY is co-released with GABA in other auditory nuclei within this circuit such as the nucleus angularis (Kuo et al. 2009) and the superior olivary nucleus (SON) (Coleman et al. 2011). Here we show that functional glycine receptors are ubiquitously expressed in all brainstem auditory nuclei.

Methods
Glycine receptor expression was characterized using immunocytochemistry and whole cell voltage clamp recordings. We used pressure application of glycine to test the functionality of glycine receptors in the NM, NL and SON. We also used pressure application and presynaptic fiber stimulation to explore the effects of glycine receptor activation on synaptically evoked inhibitory postsynaptic currents.

Results
We show that GLY receptors are present at all four of the above-mentioned nuclei throughout development of the auditory system and demonstrate that exogenous GLY application evokes strychnine sensitive currents in these neurons. Indeed, these currents were observed in nuclei such as the NM and NL, where GLY responses could not be synaptically evoked via electrical stimulation. To assess the functional role of GLY signaling we evaluated the effect of an exogenous GLY pre-pulse on synaptically evoked GABA or mixed GABA/GLY inhibitory currents in the NM and the SON, respectively. Surprisingly, activation of GLY receptors reduced the amplitude of inhibitory postsynaptic currents evoked during a 100Hz train stimulus in both nuclei. This occlusion effect was blocked during application of strychnine and recovered after
washout. This modulation was not dependent on phosphatase 2B activity but did require the flux of ions through the receptor channel.

**Conclusion**
Functional GLY receptors were located in all four primary nuclei in the chicken auditory brainstem. Pre-application of glycine suppressed the amplitude of both synaptically evoked IPSCs and spontaneous IPSCs. Activation of the GLY receptor without Cl- flux was insufficient to induce the suppression. These results suggest that ionic transmission through the GLY receptor occludes inhibition via GABA_A receptors, providing a possible modulatory mechanism for inhibition in the sound localization pathway.

**Variability of Sound Source Position and Temporal Feature Representation in Across-Frequency Integrating Neurons of the Barn Owl.**

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**Background**
The acoustic system needs to code where a sound source is located and what kind of source it is. Keller and Takahashi (J Neurophysiol 84:2638 (2000)) showed that neurons of the central nucleus of the inferior colliculus (ICC) are able to code both, the “where” and the “what”. The location of a sound source was represented by the maximum firing rate in response to interaural time (ITD) and level (ILD) differences, while temporal feature of the sound were represented by the discharge pattern of the neurons.

The acoustic pathway branches at the level of the ICC to create a forebrain and a midbrain pathway (Fig. 1). Both pathways receive input from the ICC, both integrate information of different frequency channels, and the neurons of both pathways are significantly tuned to ITD. ICC neurons exhibit a high firing rate with low variability in response to ITD stimuli (Christianson and Pena, J Neurosci 26:5948 (2006)).

**Methods**
We recorded extracellularly from single neurons of the external nucleus of the inferior colliculus (ICX, midbrain) and the auditory arcopallium (AAr, forebrain). We used “de novo” generated noise and different “frozen” noise samples to examine to what degree and how precise the sound source position and the temporal features of acoustic sounds are represented in these nuclei.

**Results**
While firing rate decreased from ICC to ICX, variability increased (Fig. 2). Preliminary data of AAr neurons showed the same effect. We could not observe any difference between the firing rate and the variability in response to “de novo” and “frozen” stimuli neither at the population (Fig. 3) nor at the neuron level.

**Conclusion**
Across-frequency integration leads to increased variability and is accompanied by a decrease in firing rate. The missing difference between “de novo” and “frozen” noise indicated a stimulus independent rate code for ITD representation in these neurons.
Previous stimulation does not affect the representation of sound source location by units of the inferior colliculus of the Mongolian gerbil
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Background
Previous stimulation, particularly the number of stimuli preceding a target stimulus has been shown to affect the localization ability (e.g., Feinkohl & Klump 2012). Here we compare the representation of sound source location in neurons of the inferior colliculus (IC) of anesthetized Mongolian gerbils in relation to the number and location of preceding stimuli with behavioral data collected from gerbils trained in an absolute and relative localization task.

Methods
In a first set of experiments, extracellular recordings from IC units in anesthetized gerbils were obtained in response to 125-ms broadband noise bursts or tones. Stimuli were presented over headphones using virtual acoustic space techniques to mimic free-field conditions. Target signals were presented from positions to the left or the right of the midline following a variable number of stimuli from the midline. Response rates and response latencies were subjected to a receiver operating characteristic (ROC) analysis. In a second set of experiments, two Mongolian gerbils were trained in a two-alternative forced choice task to indicate the direction of a target sound after the presentation of sounds from the midline that varied in number using the same stimulus types. The animals received a food reward if they approached the side of the target stimulus. Thresholds were determined applying signal detection theory.

Results
Units in the IC were spatially tuned to sounds as indicated by their response rate and/or response latency. In the vast majority of units the number of preceding sounds did not affect the spatial tuning to the target sound and thus the neurons’ discrimination ability for spatial positions as shown by the ROC analysis. Contrary to this, the discrimination ability of the gerbils performing in a localization task depended on the number of sounds that preceded the target signal. The more preceding sounds were presented, the smaller the thresholds were. Thus, while units in the IC faithfully encode the location of sound sources irrespectively of the number of preceding sounds from a reference position, behavioral thresholds are affected by stimulation history.

Conclusion
The discrepancy between the neurophysiological and the behavioral data suggests that IC neurons do not represent the observed behavior. Additional factors not related to the stimulus source location per se may affect the behavioral response.
Analysis of Localization Cues in Globular Bushy Cell with a Sound Localization Model

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Background

Globular Bushy Cells (GBCs) are one of principal cells in cochlear nucleus. They receive major excitatory inputs from auditory nerve fibers (ANFs) and project their axons to the contralateral medial nucleus of the trapezoid body. GBCs are directly involved in the sound localization pathway to the lateral superior olive. We show how GBCs improve temporal cues in spike trains by combining them with a sound localization model. The localization model is sensitive both to interaural time differences and interaural level differences.

Methods

We simulated auditory nerve responses with an auditory periphery model including power-law adaptation described by Zilany et al. (2009), which we tuned to reproduce human hearing thresholds. The GBC model was a single compartment representing a soma with Hodgkin-Huxley-like channels (Rothman and Manis 2003). It received purely excitatory inputs from ANFs. Synaptic weights were fitted to reproduce firing patterns of high-sync neurons in cochlear nucleus. They achieved a synchronization index above 0.9 and entrainment of 1 for frequencies up to 600 Hz. Simulated responses from both ears were fed to a binaural cross-correlation model (Lindermann 1986). It consisted of delay lines inspired by Jeffress (1948) and additional inhibitory elements. This combination makes the model both ITD and ILD sensitive.

Results

We simulated two types of sounds: pure tones and speech. In both cases, the sound source was moving around the listener (1 rotation/s). Our results show that GBCs greatly enhance localization cues over ANFs. GBCs are known to enhance the temporal precision of their spike trains by coincidence detection. They lock on low-frequency pure tones and enhance amplitude modulations, which are present in speech sounds. Therefore, the internal representation of the sound source location in the Lindermann model is much sharper with GBCs compared with ANFs (supplemented Figure). With speech sounds this sharpening effect is particularly pronounced.

Conclusion

The analysis of spike trains derived from GBCs and ANFs using a sound localization model showed a large improvement of localization cues in GBCs. We observed improved precision of sound source localization for simple and complex sounds. The improvement was mostly due to the higher temporal precision of action potentials in GBCs. In summary, our analysis demonstrates how temporal information is processed in the localization pathway to achieve sharp localization cues.

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Simulations of Auditory Periphery to study the Effects of Sound Level and Duration on spectral cues to Localization

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Background
Changes in sound level or duration have been shown to affect localization performance for human and cats, especially for targets in elevation. An interesting finding is that, for short-duration sound, localization performance in elevation decreases at high sound levels, whereas for long duration sound, this “negative level effect” is minimal. The integration model predicts that localization of long-duration sound benefits from “multiple looks” of consecutive short-term spectral analyses of the ongoing information. In contrast, the adaptation theory predicts that the later part of the long-duration response is more sensitive to spectrum changes than the onset. Previous studies have also suggested that a broad and flat spectrum of the sound source is critical for localization in the elevation.

Methods
An AN model (Zilany et al. 2009) for the cat was used to simulate populations of AN responses to targets in elevation. The input of the model was broadband noise filtered with cat directional transfer functions (DTFs). Three sets of broadband stimuli including long-duration random noise, short-duration (5-ms) random noise, and long-duration (100-ms) repeated noise made of identical concatenated copies of the short-duration noise were tested. The correlation between the simulated AN firing rates under these stimuli and the DTFs was used as a metric to study the level and duration effects on vertical localization.

Results
The correlation was significantly higher for the 100-ms random noise than for the 5-ms noise. However, the correlation was non-monotonic with level for sound of both durations. Replacing the 100-ms random noise with the 100-ms repeated noise had a small effect on the correlation at low and high sound levels and no effect at intermediate levels. Introducing neural adaptation during the steady-state AN response showed a large effect at high levels. The largest improvement of correlation was observed when providing the model with “multiple looks” throughout the stimulus duration.

Conclusion
The flatness of sound spectrum and peripheral adaptation benefit localization only at low and high levels, whereas neural integration that relies on “multiple looks” of the spectral analysis is critical for the peak performance at intermediate levels. The release of negative level effect observed for long-duration sound cannot be explained at the periphery and therefore, is likely a result of higher-center processing. Supported by NIH DC-00116 and DC-02840 (Yin) and DC-00376.
Effects of Forward Masking on Sound Localization in Cats

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Background
Forward masking is traditionally measured as an increased signal-detection threshold as a consequence of the addition of a preceding masking sound. The present study examined the influence of forward masking on localization of free-field sound in cats, which has not been previously studied in human or animal subjects.

Methods
Gaze positions of head-unrestrained cats were monitored using a search-coil technique. The masker speaker was fixed at the front center, while the signal could come from one of 17 speaker locations in the frontal hemifield. Cats were required to saccade from a central fixation LED to the perceived source of the target, with (i.e., the forward-masking condition) or without (i.e., the control condition) the masker. The masker was a broad-band noise (50 dB SPL, 600 to 700 ms). The target signal (25 ms) was either a broadband or a narrowband noise. The level of the signal ranged from 30 to 80 dB SPL. The gap between the end of the masker and the beginning of the target was 0 or 10 ms.

Results
When the target was a short broadband noise, the presence of the forward masker reduced localization accuracy of the target at all target sound levels, along both horizontal and vertical dimensions. The detrimental effect was larger when gap = 0 ms than when gap = 10 ms. When the target was a narrowband noise, the effect of the forward masker was considerably larger for low-frequency (0.2–2 kHz) than for high-frequency (7–15 kHz) target, in both horizontal and vertical dimensions. At certain target sound levels, forward masking had no effect on localization of the high-frequency target. For all narrowband targets, localization accuracy for targets in elevation was significantly lower than for targets in azimuth, whether a forward masker was presented or not.

Conclusion
Localization of a short sound can be affected by a preceding noise masker, even when the target was well detected. Adding a gap between the masker and the target released the masking effect to a certain degree. The detrimental effect of the broadband masker acted mostly on the low-frequency component of the target, where the interaural timing cue dominates. Supported by NIH DC-02840 (Yin).
In Vivo Response Properties of Neurons from Low-Frequency Nucleus Laminaris in the Chicken

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Background
Locating the source of a sound in the environment is a critical task for any animal. One of the most important cues animals use to accomplish this is interaural time difference (ITD), the minute difference between the moment when a sound reaches the ear closest to the source and the moment it reaches the opposite. However, how different animals compute these ITDs has been the subject of much debate. Here, we present new data from the low-frequency region of nucleus laminaris (the first brainstem nucleus that processes ITDs) in the chicken, aiming for a greater overlap in frequency range with comparable mammalian data.

Methods
Recordings were obtained from the low-frequency region of nucleus laminaris in vivo, under anesthesia, from chickens aged P28 to P48. Best frequency (BF), best interaural phase, best ITD, and frequency tuning were determined using pure tones.

Results
Approximately half of our data set is comprised of single-unit recordings, which showed no systematic differences to the multi-unit and neurophonic recordings. BFs ranged from 300 to 1600 Hz, corresponding to the caudolateral part of the tonotopically organized NL. Most interestingly, best ITDs scattered symmetrically around zero within a range of up to ±500µs, i.e., nearly equal numbers of recording sites showed a preference for contra- and ipsilaterally leading sounds, respectively. This is consistent with published, preliminary data.

Conclusion
The available data for the chicken NL now suggest a symmetrical distribution of best ITDs around a value of zero at low BFs, up to approximately 1500 Hz. At higher BFs (up to the limit of 3500 Hz in the chicken) the distribution changes to a contralaterally biased representation of auditory space. Furthermore, the range of best ITDs is in good agreement with the physiologically relevant (i.e., naturally heard) range for the chicken. These features are consistent with a Jeffress-type neural code of ITD. However, the symmetrical distribution of best ITDs around the acoustic midline at low frequencies is unorthodox. It remains to be shown whether the topographic representation of ITD, previously shown for the majority of chicken NL, also persists into the caudolateral, low-frequency region.
Mechanisms of Auditory Spatial Selectivity in Barn Owls
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Background
The physiological mechanisms of interaural time difference (ITD) encoding in birds and mammals are a matter of ongoing debate. The avian encoding strategy is assumed to be based on coincidence detection of excitatory inputs fed by neuronal delay lines, while the mammalian strategy is assumed to involve precisely timed interaction of excitatory and inhibitory inputs resulting in ITD-dependent activation balance between the left and right hemisphere (e.g., Grothe 2003, Nature Neuroscience). Given such differences it is interesting to test whether one model can explain spatial perception of sound sources both in birds and mammals. Wiegrebe and Binetti (2010, ARO) presented an experimental paradigm aimed at determining spatial selectivity in humans in a binaural unmasking task. We adopted this paradigm using barn owls as an avian species. The observed data from both humans and owls were compared to predictions of a computational model.

Methods
Three barn owls (tyto alba) were trained in a Go/NoGo paradigm using operant conditioning to detect a narrow-band noise target in broad-band masking noise. The stimuli were presented under free-field conditions, with two independent masker sources placed symmetrically to the target source in the horizontal plane. The masker was played continuously and the target duration was 110 ms including ramps. Detection threshold levels were measured as a function of target-masker separation (7.5°-110°), target center frequency (2, 8 kHz), and target location (front, left, right). The computational model is based on the equalization-cancellation model by Durlach (1963, JASA).

Results
In barn owls, the detection thresholds generally decreased with increasing target-masker separation, and there were significant interactions between both target frequency and target location with target-masker separation. The lowest thresholds were observed for an 8 kHz target in front of the subject. In humans (Wiegrebe and Binetti 2010), the thresholds generally also decreased with increasing target-masker separation and increasing target frequency, but in contrast to the owl data a target on the interaural axis leads to lower thresholds than a target in front. Especially at 8 kHz the barn owl's spatial resolution revealed by the unmasking was far superior to that of humans.

Conclusion
One computational model can explain the data in both humans and owls. Only the head-related transfer functions and the phase locking cut-off frequency need to be adjusted in the model in order to explain the observed detection threshold differences across species.

Optimizing Parameters for Efficient Acquisition of the Binaural Interaction Component of the Auditory Brainstem Response
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Background
The sound-evoked binaural interaction component (BIC) is the residual auditory brainstem response (ABR) after subtracting the sum of both monaurally-evoked ABRS from the binaurally-evoked ABR. The β peak (i.e., DN1) is the first negative peak of the BIC. Latencies of β in human and animal studies indicate a brainstem origin for this electrophysiological potential related specifically to binaural processing at the inferior colliculus. The BIC may be diagnostically important: altered β latencies and amplitudes in children and adults correlate with and predict long-term behavioral deficits in binaural processing associated with chronic conductive hearing loss. In humans, cats, chinchillas and guinea pigs, β amplitude also depends systematically on binaural cues to location, exhibiting maximal amplitude for interaural time (ITD) and level (ILD) differences of zero (midline sources); β is often no longer detectable once ITDs and/or ILDs exceed physiological range. In this regard, the BIC is also informative for perception, as changes in β peak latencies and amplitudes occurring with varied stimulus ITD or ILD are correlated with psychophysical performance (lateralization, discrimination, binaural masking level differences) in normal and hearing-impaired subjects.

Methods
While the BIC β peak is theoretically clinically useful, it can be difficult and time-consuming to measure. The ABRS from which the differential BIC is calculated are themselves very small potentials (~1 µV) measured using signal averaging to
cumulatively amplify the small potentials (such as the \( \beta \) peak). The large series of stimulus trials required for ABR acquisition typically ranges from the hundreds to thousands or more, while acquisition of a subtractive potential like the BIC \( \beta \) peak is yet more elusive. Manipulation of stimulus ITD and ILD requires even more time.

**Results**

While presentation of thousands of stimulus iterations is laborious but feasible in a laboratory setting, introduction of the BIC \( \beta \) peak as a practical clinical assessment tool is more challenging. Laboratory experiments can allow for many hours of stimulus repetitions and recordings, but such an acquisition protocol is clinically impractical. Likewise in longitudinal animal experimentation, an expedited measurement of the BIC \( \beta \) peak is also desirable.

**Conclusion**

To this end, we assess and discuss parameters for quickly and accurately obtaining reliable BIC \( \beta \) peak measurements in chinchillas. The resultant protocol allows for reliable BIC \( \beta \) peak measurement within a more clinically-feasible time period. Support: NIDCD R01-DG011555

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**Sensitivities to the relative phase of interaural-time-difference modulations between carrier frequencies**

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**Background**

Evidence suggests that the human auditory system integrates information about interaural-time difference (ITD) across frequencies (e.g., Buell & Hafter, JASA, 1991; Hill & Darwin, JASA, 1996). The present study examined the dynamic properties of the across-frequency integration mechanism, specifically the extent to which the information about the direction of ITD change is integrated across frequencies.

**Methods**

The stimulus was a 1000-ms-long complex tone (flanked with 50-ms raised-cosine onset/offset ramps), consisting of two sinusoidal carriers at 400 and 700 Hz. A sinusoidal modulation in ITD was imposed on one carrier alone or the two carriers simultaneously. The ITD of each carrier was centered at 0 \( \mu \)s, and the modulation started and ended with the zero phase. Modulation rates of 1, 2, 4, and 8 Hz were tested. ITD modulations, when imposed on the two carriers simultaneously, were in-phase or anti-phase between the carriers. Listeners with normal hearing participated in the experiments. Experiment 1 measured the threshold modulation depth for detecting the modulation with an adaptive method. The three-interval, three-alternative forced-choice (3I-3AFC) method was used; one “signal” interval contained a modulated stimulus, and the other two “non-signal” intervals contained unmodulated stimuli. Experiment 2 measured the discriminability between in-phase and anti-phase modulations. Modulation depth was fixed at a supra-threshold value (600 \( \mu \)s). In the 3I-3AFC task, the “signal” interval contained anti-phase modulations and the other two “non-signal” intervals contained stimuli with in-phase modulations. The method of constant stimuli was used to obtain the percentage of correct answers.

**Results**

Experiment 1 showed generally lower thresholds when the two carriers were modulated than when only one was modulated, indicating across-frequency integration of the information about the presence of modulation. The threshold, however, was not significantly different between the in-phase and anti-phase conditions, even when the modulation rate was as low as 1 Hz. In Experiment 2, the discrimination performance was generally at the chance level for the majority of listeners even for a 1-Hz rate.

**Conclusion**

The study failed to present compelling evidence that the auditory system is sensitive to the relative phase of ITD modulations for the conditions tested. This suggests that the directional information of even slow (~ 1 Hz) ITD modulation is lost at or below the level of across-frequency processing, or is not effectively combined at that level.
The influence of low frequency distortion products on ITD sensitivity to transposed tones
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Background
A long-standing controversy concerns brain mechanisms underlying the perception of the pitch of complex sounds, particularly the so-called ‘pitch of the missing fundamental’ in which the perceived pitch corresponds to the reciprocal of the harmonic spacing. One potential contributing factor to missing fundamental pitches is intermodulation distortion; the non-linear mechanics of the cochlea generate an excitation pattern at the low-frequency location of the basilar membrane corresponding to the periodicity in the stimulus. The salience of distortion-generated missing fundamentals (and the level of distortion generated in the cochlea) is modulated by factors such as the phase, number and order of harmonic components, the relative timing of click trains eliciting pitch percepts, and the overall stimulus level. Pressnitzer and Patterson [Pressnitzer D and Patterson RD 2001, Distortion products and the perceived pitch of harmonic complex tones. In: Breebaart, DJ, Houtsma, AJM, Kohlrausch, A, Prijs, VF, and Schoonoven, R (Eds) Physiological and Psychophysical Bases of Auditory Function, pages 97-104. Shaker Publishing BV, Maastricht, The Netherlands.] assessed the level of the distortion spectrum using the “cancellation of beats” method. They showed that adding low pass noise reduced the pitch percept for bandpass complex tones at high frequencies. Here, we assess the magnitude of intermodulation distortion using a binaural discrimination task.

Methods
We assessed the sensitivity of human listeners to ITDs conveyed in the envelopes of transposed tones as a function of sound level of the transposed tone and as a function of low-frequency noise level. ITD discrimination was determined for transposed tones with a fixed envelope ITD. Psychometric functions d’ as a function of both stimulus level and noise level were measured with a constant stimulus procedure in a forced choice paradigm.

Results
Preliminary data indicate that, in the absence of low-pass masking noise, subjects’ performance can be as good (low) as those for pure tones. Adding low-pass masking noise considerably elevates thresholds to the point that thresholds are unobtainable.

Conclusion
The exquisite sensitivity of the early auditory pathways to low-frequency binaural temporal information may mean that distortion products are particularly relevant in binaural experiments. Here we determine the minimum level of noise necessary to mask distortion products at each stimulus level.
Electrical Stimulation of the Dorsal Inferior Colliculus Modulates the Auditory Lemniscal Pathway: Implications for Tinnitus

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Background
Hyperactivity and increased neural synchrony have been linked to tinnitus. These neural changes have been observed in the central nucleus of the inferior colliculus (ICC) of tinnitus animals and patients. Thus the inferior colliculus (IC) is a potential target for treating tinnitus. We are planning a clinical trial to investigate a new IC-based auditory midbrain implant (AMI) for hearing restoration. Many of the selected AMI patients will also have tinnitus, providing a unique opportunity to assess IC stimulation effects on tinnitus suppression directly in humans. Since the AMI array can only stimulate a few IC regions, prior to implantation it is critical to determine optimal regions for tinnitus suppression. The dorsal region of the IC (ICD) is known to modulate the ICC. Therefore, our goal is to identify regions within ICD that may lead to more effective tinnitus suppression to guide AMI implantation in humans.

Methods
Multi-site electrode arrays were positioned across ICC and ICD in ketamine-anesthetized guinea pigs. Spike activity in ICC was recorded in response to broadband noise stimulation both before and after repeated electrical stimulation of the ICD to identify any residual changes in neural activity. Histological steps were taken to produce 3D reconstructions of the midbrain and to identify the location of the electrode sites across the IC.

Results
Our results reveal complex suppression and enhancement patterns. Every ICD stimulation location resulted in suppression or enhancement of at least one ICC recording location, and every recording location was suppressed or enhanced by at least one stimulation location. Interestingly, we observed a spatial gradient of suppression strength for stimulation across the ICD that could affect just one site up to all the sites across the ICC array. The caudal-medial ICD region induced the strongest suppression across the ICC.

Conclusion
Based on our data, stimulation of any ICD location causes some suppression or enhancement within the ICC, and stimulation of specific ICD regions will induce dramatic suppression across the ICC. This complex activation pattern is favorable for tinnitus in which neural changes are highly variable across patients, and thus the location of stimulation can be altered per tinnitus case. Future studies will be performed in animal models of tinnitus to verify that ICD stimulation can suppress or enhance the appropriate neural patterns to eliminate tinnitus, including suppression of spontaneous hyperactivity and hypersynchrony.

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Sound-triggered Suppression of Neuronal Firing as an Underlying Mechanism of the Residual Inhibition of Tinnitus
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Background
Tinnitus can be suppressed briefly following the offset of an external sound. This phenomenon, termed “residual inhibition,” has been known for almost four decades, although its underlying cellular mechanism(s) remains unknown. In our previous work we demonstrated that many neurons in the inferior colliculus exhibit long lasting suppression of spontaneous activity following the offset of an external sound. Hyperactivity or elevated spontaneous activity in the central auditory system often correlate with tinnitus; thus we hypothesize that sound evoked suppression underlies residual inhibition. To test this hypothesis, we studied the sound evoked suppression phenomenon in animals with behavioral evidence of tinnitus.

Methods
Experiments were conducted on CBA/CaJ mice. For tinnitus induction mice were exposed to a narrowband noise centered at 12.5 kHz presented at 116 dB SPL unilaterally for 1 hour under general anesthesia (Ketamine/Xylazine). Tinnitus was assessed with gap-induced prepulse inhibition of the acoustic startle reflex. Extracellular recordings were performed in IC both contra- and ipsilateral to the exposed ear in awake restrained animals. Pure tones at neurons’ characteristic frequency and/or wideband noise stimuli with 5 sec duration were delivered in the free-field.

Results
We found that mice with behavioral signs of tinnitus exhibited abnormally high spontaneous activity in the contralateral IC. Although most of these neurons showed abnormally high spontaneous firing, less than 25% of them exhibited sound triggered suppression of spontaneous firing. In contrast, more than 80% of ipsilateral (“unexposed”) IC neurons exhibited long lasting suppression following the offset of a sound stimulus. Bursting patterns within spontaneous firing were not evident in either the contra- or ipsilateral IC. It is important to note that even in response to a relatively short sound duration in about 10% of recorded neurons the duration of this suppression approximated the duration of the residual inhibition of tinnitus observed in humans.

Conclusion
Unilateral tinnitus induction causes a dramatic increase of spontaneous activity in the contralateral (“exposed”) IC, whereas the ipsilateral (“unexposed”) IC remains similar to the control (not exposed) animals. The majority of neurons in the contralateral “exposed” IC did not exhibit long lasting sound evoked suppression. The time course of suppression in the ipsilateral IC corresponds to the time course of residual inhibition in tinnitus patients.
This research was supported by grant R01 DC011330 from the National Institute on Deafness and Other Communication Disorders of the U.S. Public Health Service.
Background
High-pressure blast shock waves are known to cause tinnitus. Although the underlying mechanism may be due to damage to structures in the ear and/or direct brain impact that triggers a cascade of neuroplastic changes in both auditory and non-auditory centers, there is a lack of investigation of electrophysiological changes following blast-induced tinnitus. In this study, we used a rat model and investigated how blast influenced behavioral evidence of tinnitus and spontaneous firing rates (SFR) in the dorsal cochlear nucleus (DCN), inferior colliculus (IC) and auditory cortex (AC) at different time points after blast exposure.

Methods
Forty-two adult Sprague-Dawley rats were divided into three groups: one day, one month and three months after a single blast exposure in the left ear. Each group had nine blasted and five age-matched control rats. Gap detection acoustic startle reflex behavior was used for evaluating tinnitus and auditory brainstem responses were measured before and after the blast exposure. SFRs were simultaneously recorded in the DCN, IC and AC after blast exposure. At the end of experiments, all animals were euthanized and their brains were processed histologically to verify the electrode positions.

Results
Our behavior data showed that blast-induced tinnitus occurred at all frequency bands immediately after, at high frequency bands one month after, and eventually at a high- and a low-frequency band three months after the blast exposure. Compared to control rats, an early onset increase in SFR was found in the DCN of rats with tinnitus, which lasted for a month before returning to a normal level three months after the blast exposure. An increased SFR in the IC of rats with tinnitus persisted through all recording sessions. In the AC of rats with tinnitus, onset decrease in SFR occurred, which lasted for one month before reversing to an increase in SFR at three months after the blast exposure.

Conclusion
These results demonstrated that blast-induced tinnitus does not involve a uniform manifestation of increased SFR along the auditory axis. Instead, changes in SFR happen in different directions at different levels of the central auditory system. The data also suggest that blast-induced early onset tinnitus may be related to increased SFR in the brainstem whereas delayed tinnitus may be attributed to increased SFR in the IC and AC. This information may be important for developing treatment strategies of blast-induced tinnitus based on hyperactivity change patterns along the auditory axis.
Behavioural, Neural and Histological Correlates of Tinnitus in the Guinea Pig
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Background
Animal models of tinnitus are essential to furthering our understanding of neural changes accompanying the condition. Deficits in gap detection in behavioural tests and increased spontaneous neuronal firing rates (SFRs) in the inferior colliculus (IC) are used as markers for tinnitus.

Methods
We sought a histochemical marker of tinnitus in guinea pigs (GPs) by studying changes in the expression of nitric oxide synthase (NOS), a pathological marker, in the ventral cochlear nucleus (VCN). 16 GPs were subjected to unilateral noise trauma, tested behaviourally over an 8-week period and IC recordings were performed. Hearing status was assessed using auditory brainstem responses.

Results
Using strict criteria, 45% of GPs exhibited behavioural evidence of tinnitus. These same animals also showed a 30% increase in the number of NOS containing neurones in the VCN of the deafened side at eight weeks after noise exposure. By contrast, deafened animals with no behavioural evidence of tinnitus at eight weeks showed symmetric NOS staining. Increased SFRs were evident in noise-exposed GPs, when compared with control GPs, whether or not there was behavioural evidence of tinnitus.

Conclusion
The behavioural test reliably indicated the presence of tinnitus. However, our data indicate that elevated SFRs in the IC are not an unequivocal marker for tinnitus, as increased SFRs were apparent after noise exposure regardless of whether tinnitus was present or not. However, sustained changes in NOS expression in the VCN do appear to be a robust marker for tinnitus.
Unexpected increases in ipsilateral inferior colliculus spontaneous activity in a rat model of tinnitus

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Background
Although common, tinnitus is not yet well understood. One hypothesis is that damage to the peripheral auditory system changes the balance of excitatory and inhibitory inputs to the central auditory system, causing an increase in spontaneous activity (SA) in central auditory structures that is perceived as tinnitus. We measured changes in SA in the central auditory system by utilizing radioactively labeled 2-deoxyglucose (2DG). 2DG is used to measure metabolic activity in the nervous system, and we used it to evaluate changes in SA of the inferior colliculus (IC) following sound damage.

Methods
We exposed anesthetized male Long-Evans rats to a unilateral 118 dB SPL, 16 kHz pure tone for 4 hours. A minimum of two weeks after sound damage, following electrophysiological recordings, we injected rats with 2DG. Because many tinnitus studies evaluate SA in anesthetized animals, half of the animals were anesthetized with a ketamine-xylazine mix prior to 2DG injection, allowing for a direct comparison of activity in anesthetized and unanesthetized animals. Each rat was placed in a sound proof chamber for 45 minutes following injection, to allow for optimal 2DG uptake. We then sacrificed the animals, harvested the brains, and sectioned them in 40 µm sections to be placed on film. One week later, the film was developed and the radioactivity was measured in ten sectors along the tonotopic axis of the central nucleus of the IC (ICc).

Results
All four groups (control, anesthetized, n=5; damaged, anesthetized, n=4; control, unanesthetized, n=9; damaged, unanesthetized, n=15) displayed an increase in uptake along the tonotopic map of the ICc, which corresponds with our previously published data. However, the overall level of 2DG uptake was lower in the anesthetized groups compared to the unanesthetized groups. In both the anesthetized and unanesthetized damaged animals, the ICc contralateral to the damaged ear, which receives the majority of its input from that ear, displayed levels of uptake similar to that in control animals. However, the ipsilateral ICc, in both anesthetized and unanesthetized damaged animals animals, had significantly higher uptake compared to contralateral ICc or ICc in control animals.

Conclusion
The use of anesthesia appears to have an effect on the overall level of metabolic activity in the IC. Interestingly, both the anesthetized and unanesthetized groups displayed a significant increase in activity in the IC ipsilateral to the damaged ear.
Hyperactivity in Animals with a Tonal Tinnitus Percept is Narrowband

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Background
In previous work, little attention has been paid to the differences between neural responses to a sound exposure that did or did not produce a tinnitus percept. Here, neurons in rat central nucleus of the inferior colliculus were studied in noise-exposed rats characterized for the presence of a tonal tinnitus percept using a gap inhibition acoustic startle response paradigm. Recordings were done in three groups of animals: normal, exposed with tinnitus, and exposed without tinnitus. Presumably the neural correlates of tinnitus can be isolated from the effects of sound exposure in this way.

Methods
Rats were unilaterally exposed to a 16kHz tone at 120 dBSPL for 2 hours. One month post-exposure, animals were screened for tinnitus using GIASR. For electrophysiology, animals were anesthetized (im Ketamine-Acempromazine,30:1). A fenestration was made anterior to lambda and single neurons were recorded by advancing a tungsten microelectrode (AM Systems) through occipital cortex to ICC. The ear contralateral to the exposure was stimulated using a free-field speaker with tones (1 to 50kHz). Once a neuron was isolated, response maps, rate-level functions and a 10-minute recording of spontaneous activity were recorded.

Results
Relative to normal baselines, sound-exposed rats showed an increase in spontaneous activity across all BFs (Figure 1a). When sound-exposed rats were divided into tinnitus versus no-tinnitus subgroups based on behavioral screening results, the no-tinnitus group showed significant hyperactivity, while the tinnitus subgroup did not (Figure 1b). We attribute this result to the narrowband elevation of spontaneous rates in the tinnitus subgroup (Figure 2a) versus the global elevation of spontaneous rates in the no-tinnitus subgroup (Figure 3b). Within this narrowband region of hyperactivity, the spontaneous rates of the tinnitus subgroup matched the global rate elevations of the no-tinnitus subgroup (Figure 3a). Outside the region of hyperactivity, the tinnitus subgroup showed lower spontaneous rates than normal controls (Figure 3b).

Conclusion
Rats that test positive for tonal tinnitus display elevated spontaneous rates that are confined to a narrow range of frequencies. The region of hyperactivity is predicted by the tinnitus pitch, and it may be enhanced by abnormally low levels of activity at surrounding frequencies. Global hyperactivity may fail to produce a positive tonal tinnitus test either because these subjects experience a broadband tinnitus percept not captured by our current behavioral paradigm, or tinnitus is derived from the frequency-organized contrast of hyperactive and hypoactive neurons.
Co-activation of the Somatosensory and Auditory Pathways to Induce Central Auditory Plasticity as a New Approach for Treating Tinnitus

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Background
We propose a new noninvasive stimulation approach for treating tinnitus, which we call Multi-modal Stimulation Therapy (MST). An underappreciated organization of the brain for treating neurological disorders is the dense and topographic interconnectivity among the sensory, motor, cognitive, and limbic centers. Relevant for MST, receptive fields across the entire body project onto and can modulate auditory neurons. Considering that tinnitus has been associated with abnormal spiking patterns of central auditory neurons, including those within the central nucleus of the inferior colliculus (ICC) and primary auditory cortex (A1), we conceived of the idea of electrically stimulating across the body to modulate tinnitus-affected neurons. We present our initial feasibility studies in guinea pigs.

Methods
We positioned 32-site electrode arrays in different locations throughout the ICC and A1 in ketamine-anesthetized guinea pigs and recorded spontaneous and acoustic-driven activity. Acoustic stimuli consisted of pure tones and broadband noise. We compared the ICC and A1 neural activity before and after MST, which consisted of body stimulation using subcutaneous needle electrodes. For the MST paradigm, we repeatedly stimulated (≥100 trials) each of eight body regions (left leg, right leg, back, genital area, left shoulder, right shoulder, neck, tongue) paired with acoustic stimulation to induce long-term neural changes within ICC and A1. The delays between electrical and acoustic stimulation as well as stimulus levels varied across body locations.

Results
MST induced long-term changes (e.g., tens of minutes) across ICC and A1. These changes included increases or decreases in spontaneous and/or acoustic-driven activity. Interestingly, different body locations affected different central auditory neurons, which suggests that different MST parameters may be able to target specific and different neurons driving the tinnitus. Only a small percentage of auditory neurons were affected by each body location, and thus additional MST parameters (e.g., multi-site stimulation, varying pulse trains, customized acoustic stimuli) need to be investigated to induce greater auditory plasticity.

Conclusion
Our preliminary animal studies demonstrate the potential for modulating central auditory neurons using MST. Further studies need to be performed in tinnitus animals to demonstrate that MST can fix abnormal tinnitus-related neural activity, including suppression of hyperactivity and hypersynchrony. Additional body locations and stimuli will be investigated to identify effective MST parameters that can be tested in tinnitus patients.

Discovery of Somatotopy within the Inferior Colliculus: Implications for a New Tinnitus Treatment

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Background
The brain has an immense capacity to integrate and coordinate information across multiple sensory and motor modalities necessary for daily function and survival. Even a simple reflex orienting the body and head towards a visual and/or
auditory cue requires precise and rapid coordination among sensory and motor circuits. Previous studies have identified superimposed topographic maps of visual space, sound localization, and somatosensory representation within the superior colliculus. Considering the large number of somatosensory projections to the inferior colliculus (IC) and the involvement of the IC in the orienting reflex, we performed experiments to identify a similar topographic map of somatosensory responses within the IC.

Methods
We positioned a 32-site electrode array into the IC of anesthetized guinea pigs at various locations to form a 3-D grid of placements across the entire IC. We characterized the acoustic-driven responses on each site to confirm their placement within different IC subregions. We then electrically stimulated each of eight different body locations and recorded the corresponding neural activity across the IC. Histological reconstruction of the IC and the locations of recording sites were obtained for analysis.

Results
Electrical stimulation of each body site elicited excitatory responses within specific regions of the IC in a spatially-ordered pattern. Stimulation of the legs and back activated more caudal-medial regions of the IC while stimulation of regions closer to the head activated more lateral regions. Although each body region projected predominantly to a unique IC area, there were also projections to overlapping regions across IC. Additionally, PSTH shape, activation threshold and latency generally varied for different IC locations and for different body stimulation locations.

Conclusion
Our discovery of a somatotopic map within an auditory-dominant nucleus reveals an even stronger coupling and organization among the different sensory modalities than previously reported. Considering the large and well-organized projection of the body to the auditory pathways and the ability to electrically stimulate the body regions noninvasively, we conceived of the idea of activating the somatosensory pathways to modulate auditory neurons for tinnitus treatment. We are currently investigating if stimulation across the body in a coordinated pattern can induce long-term changes within the central auditory system to suppress/fix the tinnitus-affected neurons. Further details are presented in our companion poster (Benjamin Smith et al., ARO 2013).

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Generation of Induced Neurons from Cochlear Non-Sensory Epithelial Cells in Adult Mice
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Background
Spiral ganglion neurons (SGNs), which are auditory primary neurons, transmit electrical stimulation from cochlear hair cells to the cochlear nucleus in the brainstem. Once lost, SGNs do no regenerate; therefore, the development of methodologies that could be used to induce the regeneration of SGNs in a damaged ear has the potential to significantly increase the likelihood of positive outcomes for hearing impaired individuals. A recent study has demonstrated that a combination of transcriptional factors, including Ascl1, a neurogenic bHLH transcription factor, sufficed to convert mouse embryonic and postnatal fibroblasts into functional induced neurons (iNs) in vitro (Vierbuchen et al., 2010). We previously generated iNs by overexpression of Ascl1 from cochlear non-sensory epithelial cells (NSECs) at high efficiency at early developmental stages (Weichert et al., 35th ARO). In this study, our aim was to examine whether we could generate iNs from NSECs in juvenile or adult stages and to elucidate the mechanism of neural induction.

Methods
Cochleae from ICR mice at embryonic day 13.5, postnatal day 1, 5, 10 and 20 were dissected out. Following dissection and exposure of the sensory epithelium, individual cochlear explants were transfected with the Ascl1.nucEGFP expression vector or control vector. Explants were cultured for seven days in vitro and analyzed for immunohistochemistry. To determine the efficiency of induction of a neuronal fate, the percentage of transfected cells that also expressed the neuronal marker βIII-tubulin was determined for cells transfected with Ascl1.nucEGFP or a control vector pCIG.nucEGFP. Explants at E13.5 were exposed to BrdU three days after electroporation until day 7 to determine whether iNs from NSECs proliferate during conversion.

Results
91% of cells transfected with Ascl1.nucEGFP were positive for βIII-tubulin at E13.5, and this efficiency remained high at P20 (46%). No cells transfected with the control vector were positive for βIII-tubulin indicating the specificity of the result.
βIII-tubulin positive NSECs transfected with Ascl1.nucEGFP did not uptake BrdU, suggesting that direct conversion by Ascl1 occurred without cell proliferation.

**Conclusion**
Overexpression of Ascl1 is sufficient to reprogram cochlear NSECs into neurons in adult mice. Gene therapy presents an attractive strategy for treating neurodegenerative disease and direct reprogramming of mature cells from one lineage to another constitutes an alternative approach for the generation of specific cell types. Our finding is an important step in understanding and achieving the induction of regeneration of SGNs in the mature cochlea.

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**1108**

Gross Neural Responses Shown to be from the Cochlear Apex by Ototoxic Injections

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**Background**
We recently reported that the auditory nerve overlapped waveform (ANOW) recorded from the round window provides an objective measure of low-frequency (<=700 Hz) neural thresholds (Lichtenhan et al. Ear Hear. In Press). In the present study, a pharmacological method was used to assess the origin along the cochlea of simultaneously recorded ANOW, cochlear microphonics (CM), and onset action potentials (AP) evoked by low-frequency stimuli. Compound action potentials (CAP) to high-frequency (=> 2 kHz) tone bursts aided the interpretation of the low-frequency measures.

**Methods**
Ototoxic pharmaceuticals were 150 mM KCl, 250 ng/ml tetrodotoxin (TTX), or 20 mM sodium salicylate in artificial perilymph. Injection into perilymph was with a pipette sealed into the cochlear apex. Injection at this site induces a volume flow from the injection toward the cochlear aqueduct at the base of scala tympani. This results in an increase in drug concentration progressing from apex to base. Injections were started at a rate of 50 nl/min, increased to 100 nl/min after 10 mins, and to 200 nl/min after 30 mins, to partially compensate for the increasing cross-sectional area of scala tympani along the cochlear partition.

**Results**
ANOW and AP to low-frequency (353, 500, and 707 Hz), low-level (50 dB SPL) stimuli were rapidly abolished by injections, demonstrating the predominantly apical origins of these responses. ANOW and AP responses to the higher-level stimuli (65 dB SPL) were abolished later, confirming the broader origins of high-level responses. CAP thresholds to higher frequencies were abolished even later and sequentially in low- to high-frequency order, in accordance with their well-known spatial origins. Approximately 15 minutes after the start of a TTX injection, the CM to low-frequency tones was partially reduced, suggesting a neural component to CM originating in the apical turn. About 60 minutes after a KCl injection, the CM was fully abolished, suggesting the component of the CM that remains after TTX originates predominantly from hair cells in the basal turn.

**Conclusion**
The results confirm that the ANOW response from low-frequency, low-level tones originates in the cochlear apex with little contamination by basal-turn responses, and that ANOW provides a good metric for cochlear sensitivity at low frequencies.

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**1109**

Modeling the Time-Varying and Level Dependent Effects of the Olivocochlear Reflex in Auditory Nerve Responses

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**Background**
The medial olivocochlear reflex (MOCR) has been implicated as a source of benefit for listening in noisy environments. This benefit is an anti-masking effect enabled by a reduction in auditory nerve (AN) firing to continuous background noise, resulting in increased sensitivity to a stimulus of interest. MOC neurons synapse on outer hair cells (OHCs), and inhibit basilar membrane responses though the reduction of cochlear amplifier gain. The current study implements the time-
varying, characteristic frequency (CF), and level dependent effects of the MOCR within the framework of a well-established computational model for normal and impaired hearing AN responses (Zilany and Bruce, 2006) based on physiological evidence in humans and cats.

**Methods**

A second order linear system was used to model the time-course of the efferent feedback loop, and was based on measurements in humans using stimulus-frequency otoacoustic emissions. The stimulus level dependence of the MOCR was determined using a tone-in-noise paradigm, fitting model parameters to previous data from AN fibers in decerebrate cats. The model includes the effect of efferent feedback for both the ipsilateral and contralateral pathways.

**Results**

The time-course of the model, after being fit to AN data, is shown in the supplemental figure for a tone at 8 kHz with ipsilateral (solid) and binaural broadband noise (dashed). OHC gain reduction is shown as a function of time, with increasing tone level (horizontally), and with increasing noise levels (vertically). The result is a time-varying, level-dependent OHC gain dependent on both ipsilateral and contralateral stimulus amplitude. Also included is the physiologically observed CF dependent nature of the MOC efferents, which has been shown to be correlated with the innervation density of efferent terminals. The MOCR improves detection of a tone in noise by decreasing the gain of the OHCs, consistent with AN data in cats.

**Conclusion**

The AN model presented here allows us to study the complex interactions of the MOCR within the peripheral auditory system, including the bilateral interaction of stimuli to both ears. The MOCR may be important for speech recognition in noisy situations as well as for protection from acoustic trauma. Further study of this model and its efferent feedback loop may contribute to our understanding of sensorineural hearing loss in noisy situations for hearing-impaired listeners. [Research supported by NIH(NIDCD) R01 DC008327 and T32 DC00030.]
Expression of Axon Guidance Molecules in Developing Avian Auditory Pathways
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Background
The development of connections between the ear and the brain occurs with great fidelity during early embryogenesis. To explore the mechanisms underlying this process, we investigated expression patterns of five axon guidance molecules in the avian inner ear, eighth cranial nerve (nVIII) and hindbrain from E5 – E8 when the central connections are being formed. The molecules tested were chosen to survey classes of candidate molecules and include characterized molecules with opposing effects on axon growth. Further selection included molecules with suspected involvement in cranial nerve and/or inner ear development.

Methods
We performed immunolabeling of coronally sectioned chick hindbrain together with nVIII and inner ears using antibodies against Neuropilin-1 (NPN-1), Cadherin-6 (Cad-6), Semaphorin-3A (Sema-3A), Slit-2, and Netrin-1. In some cases we counterstained with Neurofilament (NF) to identify axons, TUBB-3 to identify neurons, and/or Sox10 to identify glial lineages.

Results
We found low and overlapping expression of the respectively repellant and attractant molecules, Slit-2 and Netrin-1, from E5-E6 that disappeared by E7 when central auditory connections are being formed. Patterns consistent with nVIII axon guidance were seen with NPN-1, Cad-6, and Sema-3A. NPN-1 is highly expressed in glia along the peripheral edge of the PNS-CNS junction, and in vestibular axons. Cad-6 is expressed briefly in a gradient at the PNS-CNS transitional zone as nVIII axons enter this region. Cad-6 appears differentially expressed in prospective auditory and vestibular components of nVIII. Sema-3A is expressed in patches surrounding vestibular projections, potentially consistent with its role in channeling growing axons via repulsion cues. NPN-1, Cad-6 and Sema-3A also have unique expression profiles in the central auditory nuclei n. magnocellularis (NM) and n. laminaris (NL) when NM axons project toward NL.

Conclusion
The cell-type specificity and developmental expression patterns suggest that these molecules may guide formation of auditory circuitry during embryogenesis. These observations support a role for coordinated functions of multiple classes of molecules in the formation of these pathways.
Guidance of spiral ganglion neurites by micropatterned physical cues involves TRP channels, Ca\(^{2+}\) and cyclic nucleotide second messenger systems and RhoA/Rho-associated kinase (ROCK) signaling

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Background
We recently demonstrated that microchannels generated by photopolymerization of methacrylate polymers guide neurite growth from spiral ganglion neurons (SGNs). While the signaling mechanisms activated by chemotactive molecules (e.g. semaphorins) to direct neurite growth have been widely explored, the mechanisms by which neurons sense changes in the physical environment and transduce them into directed growth remain largely unknown. Here we used dissociated spiral and trigeminal ganglion cultures to explore the contribution of TRP channels, Ca\(^{2+}\) and cyclic nucleotides, and the downstream effectors, RhoA and ROCK, to neurite guidance by micropatterned substrates.

Methods
SKF96365, gentamicin, and RNA interference were used to inhibit TRP channels. We increased [Ca\(^{2+}\)]\(_i\) by depolarizing the cultures with elevated (25 mM and 50 mM) extracellular K\(^+\) (25K and 50K). Ryanodine or xestospongin C, which inhibit ryanodine-sensitive channels or inositol-1,4,5-trisphosphate receptors, respectively, were used to decrease Ca\(^{2+}\) release from internal stores. We applied cpt-cAMP, a cyclic AMP analog, or 8-bromo-GMP, a cyclic GMP analog, to investigate the function of cyclic nucleotide second messenger systems. We further employed a FRET-based biosensor and in situ Rho GTPase activity assay to quantify RhoA activity in growth cones. Phosphorylated (activated) ROCK was detected by immunolabeling. RhoA and ROCK inhibitors and siRNA-mediated ROCK knockdown were used to investigate the contribution of RhoA/ROCK signaling to SGN neurite guidance by micropatterned substrates.

Results
SKF96365 and gentamicin each decreased neurite alignment to micropatterns. Further, growth cones immunolabeled with anti-TRPV1 antibodies and siRNA knockdown of TRPV1 decreased alignment. 25K and 50K significantly reduced neurite alignment suggesting that chronically elevated [Ca\(^{2+}\)]\(_i\) inhibits neurite alignment. Further, ryanodine or xestospongin C also decreased neurite alignment whereas inhibition of external Ca\(^{2+}\) entry with cadmium did not affect alignment. Cpt-cAMP reduced neurite alignment while 8-bromo-GMP increased neurite alignment suggesting reciprocal antagonism between these cyclic nucleotides. Micropatterns increased RhoA activity and ROCK phosphorylation in growth cones. Inhibition of RhoA reduced neurite alignment while activation of RhoA further enhanced alignment. The ROCK inhibitors, H1152 or Y27632, each reduced neurite alignment in a dose-dependent manner. Likewise, siRNA knock-down of ROCK significantly decreased alignment to the micropatterns.

Conclusion
These results suggest a signaling network that senses and mediates growth cone responses to environmental physical cues involving TRP channels, Ca\(^{2+}\) and cyclic nucleotides as second messenger systems, and RhoA/ROCK as downstream effectors.
Excitotoxic Trauma to Cochlear Afferent Synapses Is Associated with Ca\textsuperscript{2+} Entry via Ca\textsuperscript{2+}-Permeable AMPA Receptors

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**Background**

Excitotoxic trauma causes rapid loss of synapses between inner hair cells (IHCs) and type I spiral ganglion neurons (SGNs). Noise damage to synapses appears to be a consequence of such excitotoxicity. Ca\textsuperscript{2+} entry to postsynaptic terminals via NMDA receptors has been implicated as the major cause of excitotoxicity in the brain but plays a minor role, if any, in the cochlea. Here we ask whether excitotoxicity at the IHC-SGN synapse is associated with Ca\textsuperscript{2+} entry and what is the Ca\textsuperscript{2+} entry route to the SGNs.

**Methods**

Excitotoxic trauma to IHC-SGN synapses was quantified by counting contacts between IHC- SGN axon contacts in kainate-exposed four days in vitro (DIV) organotypic cochlear cultures (Wang & Green J. Neurosci., 2011). Ca\textsuperscript{2+} imaging was done using 1DIV dissociated SGN cultures (Hegarty et al. J. Neurosci., 1997). Oregon Green-BAPTA1 was used as the Ca\textsuperscript{2+} reporter. All cultures were from postnatal day 4-6 rats.

**Results**

Two-hour exposures of cochlear cultures to 100-500 \textmu M kainate caused a dose-dependent loss of IHC- SGN contacts, with complete loss at 500 \textmu M. Cotreatment with IEM1460, a blocker of Ca\textsuperscript{2+}-permeable (GluA2-lacking) AMPA receptors (CP-AMPARs) partially reduced loss of IHC- SGN contacts in 500 \textmu M kainate. In contrast, nifedipine and isradipine, blockers of L-type voltage-gated Ca\textsuperscript{2+} channels (L-VGCCs) had no protective effect, although it did potentiate the protection of IEM1460. Kainate (100-500 \textmu M) caused an increase in intracellular Ca\textsuperscript{2+} concentration, [Ca\textsuperscript{2+}]\textsubscript{i}, IEM1460 (20-100 \textmu M) completely blocked this Kainate-induced increase in [Ca\textsuperscript{2+}]\textsubscript{i}, implying that CP-AMPARs are the Ca\textsuperscript{2+} entry route in excitotoxic trauma. In contrast, the increase in [Ca\textsuperscript{2+}]\textsubscript{i} following depolarization with 30 mM K\textsuperscript{+}, previously shown (Hegarty et al., 1997) to be due primarily to Ca\textsuperscript{2+} entry via L-VGCCs, was unaffected by IEM1460. Immunohistochemistry using isoform-specific anti-GluA antibodies suggests that GluA3 and GluA4 are the predominant GluA subunits in these postnatal rats, consistent with the presence of CP-AMPARs.

**Conclusion**

Blockade of CP-AMPARs by IEM1460 reduces excitotoxic trauma and prevents Kainate-induced Ca\textsuperscript{2+} entry in SGNs in vitro. L-VGCC blockers are not protective. Taken with previous data showing that NMDAR agonists do not contribute to excitotoxic trauma at IHC- SGN synapses, we infer that excitotoxic trauma at these synapses in postnatal rats involves Ca\textsuperscript{2+} entry via CP-AMPARs, not L-VGCCs or NMDARs. (Supported by NIH R01 DC009405 & P30 DC010362)
Trade-off between reliability of spiking and high-frequency phase-locking in the barn owl’s auditory nerve

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Background
Transduction by hair cells and coding by primary afferents are crucial stages in auditory sensory coding. Auditory receptors do not reproduce every aspect of a stimulus equally well, which in turn constrains central processing. In order to accurately encode both the identity and the fine timing of complex sounds, the neural response should be as similar as possible over multiple presentations of the same stimulus while preserving phase information. However, the response pattern of single auditory nerve fibres can be highly variable. We hypothesised that the spiking mechanism, which emerges at the primary afferent synapse with the hair cell, plays a role in the variability. The barn owl’s auditory afferents are known for their ability to phase-lock to frequencies up to 10 kHz, allowing us to explore the relationship between response variability and phase locking over a broad frequency range.

Methods
We recorded the responses of single auditory-nerve fibres in anaesthetised barn owls to repeated presentations of the same noise token and characterised their phase-locking using tones and their spectral tuning using reverse correlation to unfrozen noise.

Results
Surprisingly, the increased temporal precision of auditory-nerve fibres at high best frequencies (BF) did not correlate with repeatable spike timing for complex sounds. The reliability of spiking, quantified as signal-related power, decreased with increasing best frequency, while the temporal dispersion of phase-locked discharges also decreased. However, within fibres of similar BF, there was no correlation between reliability and temporal dispersion. A spiking model with BF-dependent parameters drawn from the data was able to reproduce the decline of response reliability with increasing BF.

Conclusion
Our data suggest that reliability of discharge and spike timing precision are not directly linked in single auditory-nerve fibres. While both measures correlated tightly with BF, we found no direct relation between them. Thus, the major source of unreliability in the auditory-nerve discharge was not temporal jitter. Rather, it appeared to be a frequent failure to spike to preferred segments of the stimulus. We propose that the inability to respond to every cycle at higher frequencies is the source of increasing frequency of spiking failures. Higher-order neurons are able to mitigate such failures through convergence of many primary inputs. Thus, the representation of phase at high frequencies may come at the expense of decreased reliability of spiking. This requires an increased total number of auditory nerve fibres to adequately represent sounds.

Functional Changes in Spiral Ganglion Neurons Following Exposure to Neurotrophins

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Background
Neurotrophins such as brain-derived neurotrophic factor (BDNF) and neurotrophin-3 (NT3) provide an effective tool for the rescue and regeneration of spiral ganglion neurons (SGNs) following sensorineural hearing loss. Applied separately, BDNF and NT3 alter the firing properties of SGNs and differentially regulate ion channel expression in vitro. The present study examines the impact of combined BDNF and NT3 application on the activity of SGNs, and the contribution of hyperpolarization-activated currents (Ih).

Methods
Dissociated SGNs from early post-natal rats (P3-7) were cultured for 1 or 2 days in vitro (DIV). Untreated SGNs were grown in Neurobasal media alone, while neurotrophin-treated (NT) cultures were supplemented with both BDNF and NT3 at 50 ng/ml or 10 ng/ml each. Whole-cell patch-clamp recordings were made in both current- and voltage-clamp modes to assess neural activity and the contribution of Ih respectively.

Results
Current-clamp recordings revealed a rapidly-adapting response during sustained depolarization for 96% of SGNs (n=117), with no difference observed between NT and untreated SGNs. Resting membrane potential was consistent between all
groups, whilst input resistance was significantly reduced in SGNs exposed to 10 ng/ml neurotrophins only. Firing threshold and latency were influenced by the presence of combined neurotrophins as well as time in vitro. In voltage-clamp, membrane hyperpolarization induced a slowly activating, non-inactivating inward current sensitive to the hyperpolarization-activated (HCN) channel blocking compound, ZD7288. The ZD7288-sensitive $I_h$ increased significantly in SGNs exposed to combined BDNF and NT3. At 2 DIV, current density was $-15.1 \pm 3.0$ pA/pF ($n=21$) in untreated SGNs, compared to $-25.9 \pm 3.2$ pA/pF ($n=25$) at 50 ng/ml NT, and $-28.5 \pm 5.3$ pA/pF ($n=16$) for NT at 10 ng/ml. The increase in $I_h$ magnitude was also accompanied by a shift in the activation curve and kinetics. SGNs treated with neurotrophins at 50 ng/ml displayed a 9 mV hyperpolarizing shift in half-activation compared to untreated SGNs, whilst NT at 10 ng/ml did not display a significant shift from either group. In addition, time constants were slower at 50 ng/ml NT compared to both untreated and 10ng/ml NT SGNs.

Conclusion
Exposure to combined BDNF and NT3 increases the activity of hyperpolarization-activated currents, but does not generate a shift in SGN firing adaptation.

Dynamic Range Adaptation in the Mouse Auditory Nerve
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Background
Dynamic range adaptation refers to the shift of a neuron’s dynamic range toward the most probable levels in a continuous, dynamic stimulus, thereby improving the precision of stimulus level coding. This phenomenon has been shown at several processing centers of the auditory pathway, including the auditory nerve (AN), the inferior colliculus, and the auditory cortex, and in different species including guinea pigs, monkeys, and cats. Here for the first time, we demonstrate this phenomenon in the AN fibers of wild-type and Bassoon knockout mice. In the knockout mice, the synaptic ribbons are not anchored to the active zone of inner hair cells.

Methods
We investigated both the amount and the time course of dynamic range adaptation by recording from AN fibers in anesthetized mice using CF-tone stimuli in which the mean sound level of the stimuli switches between two values every 3 s. During each 3-s half-cycle, the level was randomly drawn every 25 ms from a distribution containing a 12-dB wide high-probability region (HPR) imposed on a uniform background.

Results
We found that AN fibers in both wild type and Bassoon knockout mice show dynamic range shift in response to these “switching-HPR” stimuli. Adaptation of AN average firing rate occurs over a few hundreds of milliseconds, comparable to that found in the cat AN (Wen et al., J. Neurophysiol, 108: 69-82).

Conclusion
Overall, our work suggests that the adaptive processing in response to changes in sound level distribution occurs at the hair-cell/AN synapse and does not require the pre-synaptic ribbon.
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The representation of vocalizations in the mouse auditory system
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Background
One of the main functions of the auditory system is to process sounds elicited by conspecifics, therefore, it is plausible that the mechanisms underlying hearing and vocalizing have co-evolved to match each other. Mice are an increasingly important model in biomedical and auditory research that seem peculiar because there is a mismatch between the acoustic features of the communication calls that they produce and the capabilities of their auditory system to process...
these sounds. In other words, we cannot as yet, explain how mice hear each other given what we know of their auditory system.

**Methods**

To investigate the mechanisms underlying the processing of high frequency vocalizations, we recorded extracellular responses of single auditory nerve (AN) fibers and neurons in the inferior colliculus (IC) of anesthetized CBA/Ca mice using a combination of glass micropipettes and silicon array electrodes. The basic response properties of neurons at these two levels of the auditory pathway were characterized using stimuli that consisted of pure tones with frequencies between 1 and 90 kHz. In addition to tones, we recorded responses to conspecific vocalizations whose frequency components were contained within the range from 60 to 90 kHz. These sounds were carefully chosen to be representative of the vocal repertoire of the CBA/Ca strain and other strains extensively used for biomedical research.

**Results**

We found that most AN fibers exhibit characteristic frequencies (CFs) that lie between 10 and 30 kHz. In contrast, CFs higher than 40 kHz are more commonly observed in the responses of neurons in the central nucleus of the IC.

**Conclusion**

These results suggest that the representation of high frequency sounds is transformed along the ascending auditory pathway and that this transformation enhances responses to vocalization stimuli.

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**1117**

**Elevated SP/AP Ratio in the Mouse Model of Endolymphatic Hydrops**

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**Background**

An elevated Summating Potential (SP) over Action Potential (AP), or SP/AP ratio, on electrocochleography (ECoG) suggests endolymphatic hydrops (ELH) in humans. The male mouse carrying the Phex<sup>Hyp<sup>−Duk</sup></sup> (Phex<sup>HD</sup>/Y), a mutant allele of the phosphate-regulating gene Phex, develops spontaneous postnatal ELH by postnatal day P21. In animals, (ECoG) has been recorded from primates, guinea pigs and rats but never on a spontaneous ELH mouse model. A survival procedure was developed and applied to the Phex<sup>HD</sup>/Y mouse to determine whether it exhibited similar electrocochleographic changes seen in humans with ELH. Concurrent histological studies would also definitely prove that an enlarged SP/AP is associated with ELH. These studies would identify direct correlation between severity of hydrops, hearing loss and the SP/AP ratio.

**Methods**

A percutaneous recording of (ECoG) was developed and applied to Phex<sup>HD</sup>/Y mice and wild-type controls at P21. The mice were then euthanized and their cochleae harvested for histological analysis. Each hydropic cochlea was assigned a grade ranging from 0 (normal) to 4 (severely hydropic). SP/AP ratios were compared between both groups using a t-test, and a linear regression test was applied to determine correlation among hydrops grade, hearing loss and SP/AP ratio.

**Results**

42 control ears and 26 Phex<sup>HD</sup>/Y ears were successfully tested. All Phex<sup>HD</sup>/Y mice ears had ELH on histology. The Phex<sup>HD</sup>/Y mice had a significantly elevated SP/AP ratio compared to the wild-type animals (mean±SD: 0.439±0.038 and 0.278±0.010, respectively, p=0.0002). The regression test between ELH and the SP/AP ratio showed no significant correlation (p=0.978). However, hearing loss was found to correlate positively with ELH grade (p=0.0002).

**Conclusion**

(ECoG) was successfully recorded in a mouse model of ELH for the first time. As seen in human studies, the Phex<sup>HD</sup>/Y mouse demonstrated an elevated SP/AP ratio. While elevation of the SP/AP shadows ELH, it does not correlate with severity ; ELH however, correlated positively with elevated hearing thresholds. Elevated SP/AP has been suggested to be linked to ELH in the literature, but this is the first study to prove that with histologic evidence. The evidence presented in this report, combined with unique advantages of the Phex<sup>HD</sup>/Y model, provides new drivers to understand the pathophysiology linked to the development of hearing loss associated with the ELH scenario, including Ménière’s disease.
Mouse Model Of The Fluctuating Hearing Loss Associated With Enlargement Of The Vestibular Aqueduct in Pendred syndrome.
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Background
Hearing loss associated with enlargement of the vestibular aqueduct (EVA) is commonly caused by SLC26A4 mutations. The hearing loss is predominantly sensorineural, variable in severity, asymmetric or unilateral, with an onset in the first few years of life. The hearing loss often shows fluctuation and overall downward progression. We previously reported a binary transgenic mouse line, in which Sltc26a4 expression is induced by doxycycline (dox). Slc26a4 expression during E16.5 to P2 was required for the acquisition of normal hearing at 1 month of age. The purpose of the present study was to characterize the natural history of hearing loss in this model.

Methods
Dox was included in the drinking water provided to the mother from the time of conception through embryonic day 17.5. ABR thresholds were measured at 1, 2, 3, 4, 5, 6, 9 and 12 months of age. Endocochlear potential (EP) was measured in the basal turn of the cochlea. We performed quantitative real-time PCR analysis of total RNA isolated from microdissected stria vascularis. PCR primers were designed to amplify cd68, cd45 and lyzs as macrophage expression markers. Plastic-embedded cochlear tissue sections of 4μm thickness were stained with toluidine blue. Numeric data are presented at average ± sem. Statistical analyses included Pearson correlation and one-way ANOVA.

Results
ABR thresholds showed fluctuating hearing loss from 1 through 3 months of age. The EP correlated with ABR thresholds at both time points, suggesting that fluctuations of hearing are primarily associated with strial dysfunction and fluctuating EP. Quantitative real-time PCR and immunohistochemistry analyses showed a significant inverse correlation of macrophage parameters in the stria vascularis with hearing ability at 1 month of age. Kcnj10 and Kcnq1 immunoreactivity appeared normal even with severe hearing loss. ABR testing at 9-12 months of age revealed progressive overall loss of hearing. Light microscopic evaluation showed atrophy and degeneration of the stria vascularis, without EVA.

Conclusion
This mouse model closely approximates the fluctuating and progressive hearing loss observed in many human EVA patients. Macrophages proliferation in the stria vascularis and loss of the EP appear to be primary events caused by a decrease of pendrin function. EVA is a radiologic marker for, but not a cause of, hearing loss. This is a valuable animal model to explore the molecular and cellular pathogenesis of, as well as potential therapeutics for, hearing loss associated with EVA. Supported by NIH R01-DC012151.

Restoration by Salicylate of Transport Function and Anion Exchanger Activity of Missense Pendrin Mutations
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Background
In the inner ear, pendrin is detected in the stria vascularis, spiral prominence and outer sulcus. It is a member of the solute carrier 26A (SLC26A) family and known as an anion exchanger involving in the conditioning of ion concentration of the endolymphatic fluid. Its mutations are responsible for hereditary hearing loss. In Japanese, 10 missense mutations, i.e., P123S, M147V, K369E, A372V, N392Y, C565Y, S657N, S666F, T721M and H723R, have been found. We have reported that salicylate restores the localization and function of mutants of prestin, which is a member of the SLC26A family and has about 45% similarity with the amino acid sequences of pendrin. Thus, it is hypothesized that salicylate restores pendrin mutants as in the case of prestin. In this study, the effects of salicylate on the localization and anion transporter activity of the 10 mutants were analyzed in vitro.

Methods
HEK 293 cells were transfected with each expression vector with the 10 mutants. After the cells were incubated with salicylate for 12 h, for the analysis of localization, they were stained with TRITC and its fluorescence was observed using a confocal laser scanning microscope. For the analysis of anion transporter activity, they were incubated in high I⁻ buffer
containing 200 kBq/ml $^{125}$I, promoting uptake of $^{125}$I, and then incubated with high Cl$^-$ buffer, leading to the export of $^{125}$I from the cells to extracellular media by transfected pendrin. Radiolabeled iodide in the cell lysates was measured using a scintillation counter.

**Results**

Immunofluorescent staining revealed that K369E and C565Y, as well as wild-type pendrin, were transported to the plasma membrane while 8 other mutants were retained in the cytoplasm (Fig. 1). Incubation with salicylate induced the transport of 4 pendrin mutants (P123S, M147V, S657Y and H723R) from the cytoplasm to the plasma membrane (Fig. 2(a)). The remaining radioactivity in cells transfected with such 4 mutants was significantly decreased by salicylate treatment as in the case of WT pendrin, indicating that their anion exchanger activity was restored (Fig. 2(b)).

**Conclusion**

Incubation with salicylate restored the transport function and anion exchanger activity of 4 pendrin mutants (P123S, M147V, S657Y and H723R). These findings suggest that salicylate possibly contribute to development of a new method of medical treatment for sensorineural hearing loss caused by pendrin mutations.
Fig. 1. Intracellular localization of pendrin and its mutants expressed in HEK293 cells. TRITC indicates pendrin. Scale bars represent 10 μm.

Fig. 2. Effects of salicylate on the function and the inhibitory effect of the cells transfected with p.P123S, p.K369E, p.H723R and p.H723R was significantly decreased by salicylate. ** P < 0.01 vs. corresponding control (cells lacking salicylate), *** P < 0.001 vs. empty without salicylate.
A Balanced Chromosome Translocation Reveals a Predicted Lipase’s Involvement in Sensorineural Hearing Loss
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Background
Chromosomal translocations, an abnormal exchange of genetic material between chromosomes, are estimated to be as frequent as 1 in 500 newborns and may result in abnormal phenotypes due to gene disruption or dysregulation. The Developmental Genome Anatomy Project (DGAP, www.DGAP.Harvard.edu) systematically examines subjects with chromosomal translocations and abnormal phenotypes to identify disrupted genes etiologic in the disorder. Of particular interest for DGAP are chromosomal translocations associated with congenital hearing loss. DGAP056 is an archetypal case with congenital profound sensorineural hearing loss, as well as a constellation of other symptoms including mild craniofacial abnormalities (coloboma, exotropia, blepharophimosis, and low-set posteriorly rotated ears), mitral valve prolapse, hypospadias, and early onset prostate cancer.

Methods
Several standard methods were used including protein homology modeling, karyotyping, fluorescent in situ hybridization (FISH), 3’ RACE, qPCR, whole-mount in situ hybridization (WISH), mouse KO, Western blot (WB), immunohistochemistry (IHC), ABR and DPOAE.

Results
The karyotype, confirmed by FISH and PCR analysis, reveals that DGAP056 has a de novo apparently balanced translocation involving chromosomes 2 and 13, t(2;13)(p24.1;q22.3)dn. The translocation results in a breakpoint in chromosome 2 that disrupts an unannotated gene, designated C2ORF43. Analysis of the subject’s immortalized lymphoblast cells reveals a marked reduction in C2ORF43 expression, as well as expression of two abnormal, truncated isoforms. Homology analysis indicates that C2ORF43 produces an evolutionarily conserved protein that is part of the alpha/beta hydrolase clan of proteins, many of which act as serine-based lipases, whose central motif, the catalytic triad, is dismantled in the abnormal transcripts. Further investigation of C2ORF43 was carried out in model organisms. WISH demonstrated that staining of C2ORF43 orthologs was localized to the developing inner ear of both zebrafish and mice. A newly developed mouse KO model demonstrated, by WB and IHC, that protein synthesis of the C2ORF43 ortholog was essentially absent in the KO mouse ear compared to the WT. Physiological measurements of hearing, using ABR and DPOAE, indicate that at six months, some KO mice exhibit signs of hearing loss.

Conclusion
C2ORF43 appears to play an important role in development and/or maintenance of the inner ear. While it is unclear what the role of this putative serine-based lipase might play in the inner ear, analysis of DGAP056’s chromosomal translocation has revealed some of the genetic components of hearing loss and highlights the power of the DGAP model.

ACEMg Diet Supplement Influences Progression of Hereditary Deafness
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Background
Dietary supplements have been shown to reduce auditory insults caused by aminoglycosides and overstimulation. Vitamins A-C-E and Mg (ACEMg) have been shown to reduce cochlear acoustic and ototoxic trauma. When given to an 11-year-old boy with Connexin 26 (GJB2) mutations 35delG/167delT and progressive hearing loss, hearing thresholds showed an average improvement of 2 dB/year over two years. To understand the mechanism of this outcome and extend it to other forms of hereditary hearing loss, we tested the influence of ACEMg on hearing loss and pathology in two mouse models of human mutations, DIAPH3 (AUNA1) and GJB2 (DFNB1).

Methods
Gjb2 conditional knock-out (Gjb2-CKO) mice were created by breeding Gjb2loxPloxP mice with Sox10-Cre PAC mice, abolishing expression of Gjb2 in supporting cells of the organ of Corti. Diap3 (mouse DIAPH3 ortholog) transgenic mice
Viral-mediated Gjb2 Gene Expression Partially Restores the Gap junction Function in the Cochlea of Conditional Connexin26 null Mouse

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Background

Mutations in the Gjb2 gene, which codes for connexin 26 (Cx26), are the most common causes of human nonsyndromic hereditary deafness. Non-sensory supporting cells in the cochlea are connected extensively by gap junctions (GJs), heteromerically assembled from both Cx26 and Cx30, to facilitate intercellular ionic and biochemical coupling. The cCx26 null mouse model is deaf and intercellular coupling among the sensory supporting cells in the cochlea are severely impaired.

Methods

Cx26 was tagged with green fluorescence protein (GFP) and inserted into an adeno-associated viral vector driven by the CMV promoter (AAV1-CMV-Cx26-eGFP). The recombinant viruses were directly injected into the scala media of one side of the cochlea of early postnatal (either P0 or P1) cCx26 null mice in order to rescue the impaired GJ functions. The contralateral side was used as a control for the same animal. In order to check the effect of surgery procedures on the hearing, the same injections were performed in wild type mice.

Results

We found that the hearing thresholds were normal in wild type mice observed one month after injections as measured by ABR tests. The earliest Cx26-GFP expression was detected about 5 days after injections and lasted for at least 3 months. The expression was found in many types of supporting cells of the organ of Corti, fibrocytes of connective tissues and marginal cells in the stria vascularies, with the strongest Cx26-GFP signal found in the outer sulcus cells, marginal cells and spindle shaped cells. We found significantly more cells in the injected side survived in the sensory epithelium of cCx26 null mouse at one month. GJs biochemical coupling in the supporting cells was tested using the fluorescent dye transfer assay (injection of propidium iodide, PI) and the fluorescent recovery after photobleach (FRAP) assay. Both PI transfer and FRAP assays demonstrated a significant amount of restoration of GJ-mediated biochemical coupling among the outer sulcus cells in the injected cochlea of cCx26 mice. However, the tunnel of Corti remained closed and the hearing test of the injected mice did not show significant hearing recovery so far.

Conclusion

This study demonstrated partial restoration of GJ-mediated biochemical coupling in the cochlea of cCx26 null mice after gene delivery via AAV1-CMV-Cx26-eGFP. Late and incomplete expression by various types of cochlear cells may be responsible for the failure of hearing rescue in the cCx26 null mice.
Absence of tricellulin affects the ultrastructural organization of the tricellular tight junctions and leads to degeneration of cochlear hair cells

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Background
In the cochlea, tight junctions between the epithelial cells lining the scala media are responsible for isolating the endolymph from the perilymph, which is indispensable for normal cochlear function. Mutations in TRIC, which encodes a tricellular tight junction protein, are known to cause autosomal recessive nonsyndromic hearing loss (DFNB49) in humans, but the exact mechanism leading to deafness is unknown.

Methods
We generated a mouse model for DFNB49 that carries a mutation in Tric, p.R497*, orthologous to a previously identified mutation, p.R500*, in humans. Inner ear function was determined by ABR, DPOAE and vestibular evoked potential measurements. The cochlear and vestibular sensory epithelia were examined by immunofluorescence and scanning electron microscopy. The effects of the mutation on the ultrastructure of tight junctions in the inner ear were determined by freeze fracture microscopy. Tracer assays and endocochlear potential measurements were performed to establish the integrity and functional state of the stria vascularis. Whole cell patch clamp recordings were performed to determine the functional maturation in early postnatal outer hair cells (OHCs). Serum chemical, hemotological and histological analysis were performed by using the services of the Division of Veterinary Resources at NIH.

Results
Mice homozygous for the p.R497* knockin mutation displayed early onset rapidly progressing hearing loss and were completely deaf by P30. Progression of hearing loss coincided with OHC degeneration, followed by loss of inner hair cells. The mutation neither affected the maturation of voltage-gated ion channels in young postnatal OHCs, nor endocochlear potential nor vestibular evoked potentials. However, the mutation appeared to affect the ultrastructure of the tricellular tight junctions in the reticul lamina of the organ of Corti, marginal cell layer of the stria vascularis and the vestibular epithelia. Consistent with the hypothesis that the barrier function of the reticul lamina is lost in Tric knockin mice, the hair cell loss in these animals was rescued in vivo when the endocochlear potential was eliminated by deleting Pou3F4 that is necessary for normal development of the stria vascularis. Finally, the Tric knockin mice showed phenotypic changes in mandibular salivary glands, thyroid follicles, olfactory epithelium and heart.

Conclusion
Loss of tricellulin leads to organ of Corti degeneration that is likely due to defective paracellular barrier function of the reticul lamina. Although mutations in tricellulin are known to cause non-syndromic deafness in humans, orthologous mutations in mouse result not only in deafness but also in abnormalities beyond the inner ear.

Connexin26 (Cx26) deficiency impairs miRNA expression in the cochlea

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Background
Gap junctions play an important role in hearing. Connexin 26 (Cx26, GJB2) mutations are responsible for 50-70% of nonsyndromic hearing loss. Cx26 deficient mice demonstrate cochlear developmental disorders, reduction of endocochlear potential (EP), and degeneration of cochlear cells including hair cells. We recently found that cell degeneration is not a primary cause for Cx26 deficiency induced hearing loss (Liang et al., 2012). Some mechanisms other than cell degeneration, such as cochlear development disorders, may play an essential role in this prevalent genetic hearing loss. However, the molecular genetic mechanism underlying cochlear developmental disorder associated with Cx26 deficiency is little known. During organic development, intercellular communication between cells is necessary and
required for synchronizing cell proliferation and differentiation. MicroRNAs (miRNAs) are naturally-occurring regulatory RNAs and play critical roles in cell differentiation and organisms. In another report (Zhao et al., 2013 ARO meeting), we have demonstrated that miRNA can pass through gap junctions. In this study, we studied whether miRNA can pass through inner ear gap junctions in vivo and whether Cx26 deficiency can impair miRNA expression and intercellular communication in the cochlea.

**Methods**
Fluorescence-tagged miRNA intracellular injection was used to assess miRNA intercellular diffusion between the cochlear native cells. We also used quantitative real-time PCR (qRT-PCR) to measure miRNA expression in the Cx26 deficient mouse cochlea during postnatal development.

**Results**
By intracellular injection of fluorescence-tagged miRNA, we directly demonstrated that miRNA could pass through gap junctions between cochlear supporting cells. Gap junctional blockers could eliminate this miRNA intercellular transferring. Using Cx26 knockout mice, we further found that Cx26 deficiency disrupted miRNA intercellular transferring in the cochlear sensory epithelium and reduced miRNA expression in the cochlea during postnatal development.

**Conclusion**
These studies provide direct evidence for miRNA intercellular exchanges between native cochlear cells in vivo. These studies also provide important information about gap junctional function in the regulation of cochlear development.
Mutation in a novel actin-binding protein, exhibiting an Arl2 GAP activity, is associated with nonsyndromic deafness DFNB88

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Background
Hearing loss is an etiologically diverse condition that can be genetic in origin. While mutations in over 100 loci have been causally linked with hearing impairment, less than half of them have had their gene identified. The purpose of our study was to identify a new gene responsible for nonsyndromic recessively inherited prelingual hearing loss in a large human pedigree.

Methods
We have enrolled a large consanguineous Pakistani family PKDF468 segregating prelingual, recessively inherited, nonsyndromic, moderate to profound sensorineural hearing loss. Genome wide analysis was performed after excluding the linkage to known recessively inherited nonsyndromic loci (DFNB). Whole exome followed by Sanger sequencing was performed using the DNA from the affected individuals of the PKDF468 family. Fluorescently tagged constructs were used to evaluate the effect of the mutant allele. Cosedimentation and GTPase activating protein (GAP) activity assays were performed as initial characterizations of this newly identified deafness protein.

Results
We mapped a new deafness locus, designated as DFNB88 on chromosome 2p11 in family PKDF468. Our sequencing analyses revealed a missense mutation in a candidate gene encoding a novel actin-binding protein. The DFNB88 protein is largely uncharacterized with no known biochemical activities reported to date. Our RT-PCR studies demonstrated the expression of DFNB88 protein in both cochlear and vestibular epithelia of mouse. In cochlear explant, fluorescently tagged DFNB88 protein is targeted to the actin-base structures of hair cells. Similarly, in CL-4 cells, it targets to the microvilli on the cell surface while this ability is impaired in the mutant protein with missense mutation identified in family PKDF468. DFNB88 protein is also colocalized with actin-cytoskeleton in MDCK cells. When treated with cytochalasin D, MDCK cells show a disrupted actin cytoskeleton that leads to the mislocalization of the DFNB88 protein. We found a direct interaction between F-actin and DFNB88 protein in a highspeed cosedimentation assay. Moreover, DFNB88 protein possesses GAP activity against Arl2 GTPase. Arl2 belongs to the Ras superfamily of GTPases, containing multiple members known to be involved in the regulation of the actin cytoskeleton architecture and membrane traffic.

Conclusion
Taken together, these data suggest that the DFNB88 protein is linked to the actin cytoskeleton and might regulate it through small GTPases. Our data expand the genetic basis of nonsyndromic hearing loss and highlight the indispensable function of DFNB88 protein in sound perception. To our knowledge, this is the first report of this protein associating with hearing impairment in humans.
Protein Prep Kit, Expression Pathology, Rockville, MD) for celloidin-embedded temporal bones. For shotgun MS analysis, tandem 2D chromatography MS/MS using an LTQXL Orbitrap (Thermo Fischer) was employed. The MS/MS data analysis and peptide matching to FASTA human databases was done using SEQUEST and proteome discoverer software.

Results
Several proteins were revealed in both normal otic capsule bone and otosclerosis-infiltrated otic capsule, and many were identified only in otosclerosis. Of interest, TGFβ1 was identified in otosclerosis but not in the normal control temporal bone specimen. Aside from TGFβ1, many proteins and predicted cDNA-encoded proteins were observed, with implications in cell death and/or proliferation pathways, suggesting a possible role in otosclerotic bone remodeling. Immunostaining using TGFβ1 monoclonal antibody (pTyr369, Novus Biologicals, CO) directed against the extra-cellular matrix revealed marked staining of the spongiotic otosclerosis lesion and scant staining in normal bone adjacent as well as in the control temporal bone.

Conclusion
Mechanisms involved in cochlear extension of otosclerosis are still unclear, but the implication of TGFβ1 is supported by the present proteomic data. Our immunostaining results support this data. The established role of TGFβ1 in the chondrogenesis process support the theory of a reaction targeting the globulae interossei within the otic capsule.