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Noise exposure immediately activates cochlear mitogenactivated protein kinase signaling

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Abstract

Noise-induced hearing loss (NIHL) is a major public health issue worldwide. Uncovering the early molecular events associated with NIHL would reveal mechanisms leading to the hearing loss. Our aim is to investigate the immediate molecular responses after different levels of noise exposure and identify the common and distinct pathways that mediate NIHL. Previous work showed mice exposed to 116 decibels sound pressure level (dB SPL) broadband noise for 1 h had greater threshold shifts than the mice exposed to 110 dB SPL broadband noise, hence we used these two noise levels in this study. Groups of 4-8-week-old CBA/CaJ mice were exposed to no noise (control) or to broadband noise for 1 h, followed by transcriptome analysis of total cochlear RNA isolated immediately after noise exposure. Previously identified and novel genes were found in all data sets. Following exposure to noise at 116 dB SPL, the earliest responses included upregulation of 243 genes and down-regulation of 61 genes, while a similar exposure at 110 dB SPL up-regulated 155 genes and down-regulated 221 genes. Bioinformatics analysis indicated that mitogen-activated protein kinase (MAPK) signaling was the major pathway in both levels of noise exposure. Nevertheless, both qualitative and quantitative differences were noticed in some MAPK signaling genes, after exposure to different noise levels. Cacnalb, Cacnalg, and Pla2g6, related to calcium signaling were down-regulated after 110 dB SPL exposure, while the fold increase in the expression of Fos was relatively lower than what was observed after 116 dB SPL exposure. These subtle variations provide insight on the factors that may contribute to the differences in NIHL despite the activation of a common pathway.

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Keywords

Acoustic trauma; mitogen-activated protein kinase signaling; noise induced gene expression; noise induced hearing loss; permanent threshold shift

Introduction

The National Institute for Occupational Safety and Health estimates that 22 million American workers are exposed to hazardous noise, with a third of those workers demonstrating measurable noise induced hearing loss (NIHL).^[1] Two types of hearing loss have been described following noise exposure. Permanent threshold shift (PTS) is an irreversible increase in hearing threshold due to loss of or changes in cochlear sensory cells and corresponding nerve fiber degeneration. In contrast, temporary threshold shift (TTS) is a relatively short-term transient loss of hearing that can resolve to a prenoise exposure level in minutes, hours, or days. However, in both TTS and PTS, the pattern of hearing loss varies depending on the differences in the noise paradigm to which they were exposed. Elucidation of the early molecular signatures that correspond to different levels of noise is a critical step in the prevention of NIHL.

Meltser *et al.* compared changes in certain cochlear proteins following noise exposures to TTS levels (100 dB SPL for 45 min) and PTS levels (110 dB SPL for 2 h) in CBA/CaSca mice sampled at 30 min, 2 h, and 24 h postexposure.^[2] They found differences in three mitogen-activated protein kinases (MAPK) (ERK1/2, p38, and JNK1/2) between the acute phase of TTS and PTS: 24 h after exposure, the MAPK levels in TTS mice had not yet stabilized while it stabilized in PTS mice. In both exposures, brain-derived neurotrophic factor (BDNF) was elevated, but the elevation persisted only after PTS. BDNF's receptor TrkB was down-regulated only after PTS. Although these data suggest that TTS and PTS are linked to two distinct processes, the differences in the molecular responses were detected at a much later time point after noise exposure. Since early intervention is crucial to the prevention of NIHL, it would be important to evaluate the molecular responses immediately after noise exposure, especially at intensities that are capable of inducing different levels of PTS.

Protective factors against NIHL implicate three contributors to noise-induced loss and hearing damage: Reactive oxygen species (ROS), intracellular stress apoptotic pathways, and excitotoxicity due to glutamate neurotransmission.^[3] For ROS, Henderson *et al.* has hypothesized that the mechanism of PTS is dependent upon necrotic and signaled cell death generated by free-radical damage.^[4] In animal models, cochlear treatment with both exogenous and endogenous forms of anti-oxidants such as R-N6-phenylisopropyladenoisine,^[5] glutathione,^[6] N-L-acetylcysteine, creatine,^[7] tempol,^[8] and vitamin C^[9] have been reported to be protective against PTS in preexposure and postexposure regimens. Similarly, blockade of intracellular stress pathways such as c-Jun N-terminal kinase and induction of heat shock protein Hsp70 have been shown to be protective against noise-induced cell death.^[10,11] Another hypothesis is that the excessive noise leads to glutamate excitotoxicity: Caroverine (a glutamate receptor antagonist) and riluzole (a

glutamate release blocker) have also been efficacious in preventing cell death and subsequent hearing loss.^[11,12] Despite these advances, however, therapies aimed at molecular targets linked directly to different patterns of NIHL remain to be developed.

While microarray analysis can provide clues into the regulatory pathways controlling NIHL, a relative paucity of studies has evaluated global gene expression in the period immediately following noise exposures.^[13,14] Understanding the early molecular events that led to a shift in hearing threshold may provide insights toward the development of targeted therapeutic interventions against NIHL. Therefore, the purpose of this study was to look at gene expression immediately after noise exposure that induces different levels of hearing loss.

Methods

Figure 1 provides a schematic diagram of the experimental procedure used in this report.

Subjects

The study used 72 4–8-week-old female CBA/CaJ inbred mice (Jackson Laboratories, Bar Harbor, ME, USA). Animal procedures were approved by the University of Cincinnati Institutional Animal Care and Use Committee. The animals were housed, and all procedural activities were conducted in an Association for Assessment and Accreditation of Laboratory Animal Care accredited facility at the University of Cincinnati adhering to Laboratory Animal Medical Services policies and guidelines.

Auditory brainstem response evaluation

One week prior to noise exposure, the mice were tested for their ability to generate a normal auditory brainstem response (ABR) at 4, 8, 16, and 32 kHz. ABR data were recorded using Tucker-Davis Technologies System II hardware (Alachua, FL, USA) with BioSig[®] controlling software. In order to show that our exposure procedure can cause PTSs, a separate (test) batch of mice was exposed to identical conditions, and ABR recorded for the test batch of mice. The permanent effect of noise exposure on hearing is usually measured several days after noise exposure. In the test batch reported here, we recorded ABR 11 days after noise exposure. The ABR evaluation protocol described previously was used here without any change.^[15]

Exposures

The mice were randomly divided into three groups exposed to no noise (control), exposed to 110 dB SPL, or exposed 116 dB SPL broadband noise for 1 h. The acoustic system consisted of a General Radio (Concord, MA, USA) Type 1310 random noise generator whose output signal was controlled by a Mackie (Loud Technologies Inc., Woodinville, WA, USA) mixer amplifier. The output was amplified by a Soundcraftsman[®] 4×300 Power Amplifier (Santa Ana, CA, USA), a single channel of which drove four Realistic super-tweeters (Radio Shack Corp, Ft. Worth, TX, USA, #40-1310B) built into the removable noise chamber cover. A graph of the one-third octave analysis of the broadband noise has been presented elsewhere.^[16] It is virtually flat between 8 and 20 kHz. Calibration and monitoring of the noise level inside the chamber was performed by a Sennhauser MKE 2–3

electret microphone whose input was displayed and analyzed on a Brüel & Kjær 2608 Measuring Amplifier. Microphone calibrations were confirmed preexposure and postexposure by a G.R.A.S. 42AB Sound Level Calibrator (Holte, Denmark).

Tissue collection

Immediately following noise exposure, the mice were anesthetized with an overdose of tribromoethanol (Avertin). Temporal bones were immediately dissected from the skull. The cochlea was isolated and cleaned of bone and soft tissue under a dissecting microscope. RNALater[®] (Ambion Inc., Austin, TX, USA) was continuously applied over the inner ear. The cochleae were stored in 1.8 mL capped centrifuge tubes filled with RNALater[®] at ambient temperature, placed in an insulating container and shipped overnight to the Alagramam Laboratory.

Total RNA extraction

To conduct the experiment in triplicate, a total of 72 mice in three batches were used. Each batch consisted of 24 mice: 8 mice for 116 dB exposure, 8 mice for 110 dB exposure, and 8 mice for 0 dB (control). The mice exposed to noise for 1 h were sacrificed immediately, and the cochlear tissue was harvested. Cochleae from each exposure sub-group (116, 100 or 0 dB exposure) were pooled together in a ratio of 0.75 mL of Trizol (Invitrogen, Carlsbad CA, USA)/50 mg of sample. A pellet pestle (Kimble-Chase, Vineland, NJ, USA) was used to homogenize the samples. The resulting homogenate was incubated at room temperature for 5 min. Chloroform was then added to the mix at a ratio of 0.2 mL/0.75 mL of Trizol to induce phase separation. Samples were mixed by manually inverting the tube for 15 s, and incubated for 15 min at room temperature. The samples were centrifuged at 12,000g for 15 min at 4°C and the aqueous phase containing the total RNA was removed. RNA was precipitated from the aqueous phase through the addition of 0.5 mL isopropyl alcohol per 0.75 mL of Trizol. Samples were incubated at room temperature for 10 min and centrifuged at 12,000g for 10 min at 4°C. The RNA pellet was washed in 75% ethanol by vortexing and centrifuged at 7500 g for 5 min at 4°C. After a brief air-dry, the RNA was resolubilized in RNAase free water. Total RNA samples were further purified using an RNeasy Mini Kit (Oiagen Co., Chatsworth, CA, USA). Final samples from each exposure sub-group vielded between 0.75 and 1.75 µg/µl of total RNA. An aliquot of the samples was retained for further quantitative real-time polymerase chain reaction analysis, and the rest was forwarded to the Affymetrix Resource Facility at Yale University (http://ycga.yale.edu/index.aspx) for microarray hybridization.

Microarray data generation

Total RNA received by the Affymetrix Resource Facility was analyzed for quantity and purity. Absorbance ratios, measured by Nanodrop (Wilmington, DE, USA), were used to assess the amount and quality of the samples (2.1 A260/280 1.8 and A260/230 1.8). Further quality control was conducted with an Agilent 2100 Bioanalyzer (Foster City, CA, USA), which was used to score the RNA samples via an RNA Integrity Number (RIN). Measurements from the Nanodrop and Bioanalyzer (RIN >8) ensured the quality of RNA used for microarray hybridization. As described by the manufacturer, the Affymetrix Mouse

Genome 430 2.0 Array (Santa Clara, CA, USA) contains an excess of 45,000 probe sets, with >34,000 characterized genes represented. In the present study, nine 430 arrays were used with each of the three exposure groups (control, 110 dB SPL, 116 dB SPL) performed in triplicate. Each array represents the pooled contribution of eight mice. Hybridization, washing, staining, and scanning of the 430 2.0 GeneChip were performed in accordance with the protocol outlined in the Affymetrix GeneChip Expression Analysis Technical Manual ver. 5 (Santa Clara, CA, USA). Briefly, 5 µg total RNA from each exposure experiment was reversed transcribed with Oligo (dT) primers generating first strand cDNA. First strand cDNA was used to generate double-stranded cDNA, which in turn was used as a template for the production of biotin labeled complementary RNA (cRNA) in an in vitro transcription reaction. About 20 µg of biotin-labeled cRNA was fragmented and later added to a hybridization cocktail used in a 16 h incubation. Washing, staining, and scanning steps were coordinated via the GeneChip Operating Software (GCOS) program (Santa Clara, CA, USA). Wash and stain steps were performed with an Affymetrix Fluidics Station 450. Scanning was performed using the Affymetrix GeneChip 3000 7G Scanner. The converted digital intensity signals were captured as a raw image file (.dat) and further refined by GCOS (.cel files).

Microarray data analysis

Files generated by GCOS (.cel files) were imported into Partek Genomics Suite (St. Louis, MO, USA) where the GC-RMA algorithm was applied for background noise correction, normalization, and summarization of the data set into individual expression measures.^[17] Analysis of variance (ANOVA) was applied to this data set, within the Partek Genomics Suite, to identify differentially expressed probes across different exposure conditions.^[18] This analysis returned a fold-change (FC) for each probe across different treatments (110 vs. control and 116 vs. control) and a *P* value for each comparison. For the purposes of this study, a minimum 1.2-FC (up- or down-regulated) was required for a probe to be considered a candidate for differential expression. A Benjamini-Hochberg false discovery rate (FDR) analysis was applied to the candidates so as to rule-in differentially expressed genes while controlling for the number of false positives present within the data set.^[19] A 15% FDR for 110 versus control and 116 versus control was chosen. Though selection of a suitable FDR cutoff is somewhat arbitrary, an FDR of 15% was selected on the basis of ruling "in" as many previously identified noise induced genes (from comparable microarray expression studies) as possible while keeping the FDR percentage relatively low.^[13,14,20,21]

Pathway analysis

Differentially expressed genes, as determined by the Benjamini-Hochberg analysis, were entered into a gene-ontology/pathway analysis tool: DAVID [Database for Annotation, Visualization and Integrated Discovery].^[22] As of this writing, DAVID (http://david.abcc.ncifcrf.gov/) is freely available online. The enriched list from the previous microarray data analysis was entered into the program, which queried the entered list, looking for the annotations (e.g., gene ontology [GO] terms, pathways) that were enriched in comparison to what would be expected relatively to the background. The background was set to the entire genome-wide genes relevant to *Mus musculus*. DAVID's functional annotation chart module uses a modified Fisher's exact test (EASE score) to calculate gene-

enrichment for a particular annotation.^[23] An EASE score is modified in the sense that it removes one gene from a category of interest and calculates Fisher's exact test with the remaining queried genes. By removing one gene, annotations with few gene members are

greatly penalized, while slightly handicapping more robust annotations. This conservative approach allows ruling-out of less significant annotations due to false positive members. A smaller EASE score indicates greater enrichment, with a cutoff of 0.05 indicating significant enrichment.

Quantitative real-time polymerase chain reaction

Following manufacturer instructions, cDNA was generated by using the Superscript First-Strand Synthesis System for quantitative real-time polymerase chain reaction or qPCR (Invitrogen, Carlsbad, CA, USA) on the total RNA recovered from each of the previous exposures. About 500 ng of RNA pooled from each exposure group were used in separate reverse transcription reactions. Oligo-(dT)₁₂₋₁₈ and random hexamer primers were used simultaneously in each reaction mix. The resultant 20 µl reaction mixtures were diluted 10fold using RNAase free water and were later used for qPCR. Primer3 (http:// frodo.wi.mit.edu/primer3/input.htm) was used in the design of each primer pair [Table 1]. When possible, primers were designed to span at least 1 intron, thereby allowing discrimination between cDNA amplified bands and possible genomic contamination. A final primer concentration of 200 nM of each forward and reverse primer was used per qPCR reaction. Sybr Green I (Sigma, St. Louis, MO, USA), a double-stranded-DNA binding dye, was used to dilute to a final concentration of $\times 0.4$. qPCR reactions were monitored using the DNA Engine Opticon 2 qPCR detection system (Bio-Rad Laboratories, Inc., CA, USA). Opticon Monitor software (v 3.0, Bio-Rad Laboratories, Inc., CA, USA) gathered realtime data. The PCR cycle used for each reaction was as follows:

- **1.** 2 min-94.0°C,
- **2.** 20 s-94°C,
- **3.** 20 s-55°C,
- **4.** 15 s-72°C,
- 5. Plate read,
- 6. Jump to steps 2, 3, and 4 four more times,
- 7. Melting curve 55–99°C.

All samples, including the housekeeping gene b-actin, were run in triplicate. Reactions with no cDNA template added, and reverse transcriptase-minus reactions with cDNA template provided the necessary negative controls. Specific products were verified by running products on 1.5% agarose gels with added ethidium bromide and melting curve analysis (data not shown). FC was calculated using the Pfaffl method.^[24] The main distinction between using the common 2-DDCt method versus the Pfaffl method, is that efficiencies are assumed to be equal across primer pairs in the former method while their differences are accounted for in the latter. In other words, if the efficiencies between two separate PCRs

were both 100%, the Pfaffl method would simplify to the 2-DDCt method. Efficiencies of each primer pair were calculated using the real-time PCR miner approach.^[25]

Results

In this study, 4–8-week-old CBA/CaJ mice were exposed to two different noise conditions (116 dB SPL and 110 dB SPL) for 1 h, and to no noise (0 dB or control). Before proceeding with the main experiment (described below), we used a separate batch of CBA/CaJ mice to confirm that exposure to broadband noise at 110 and 116 dB SPL for 1 h results in ABR threshold shifts. The hearing loss induced by 116 dB SPL was 10–15 dB higher than that induced by 110 dB SPL noise exposure [Figure 2]. For the main experiment, immediately following noise exposure, the whole cochleae from each exposure group were harvested, and total RNA extracted for microarray hybridization and qPCR. Both microarray hybridization studies and qPCR analysis were performed in triplicate for each exposure. ANOVA analysis of the microarray data returned a relative FC and *P* value for each comparison (116 dB SPL vs. control, and 110 dB SPL vs. control). Enriched datasets were identified on the basis of FC and Benjamini-Hochberg analysis (FDR = 0.15-0.17).

The differences in gene expression immediately following 1 h of 116 dB SPL noise exposure versus no noise exposure (116 vs. control) were compared. After applying Benjamini-Hochberg analysis (FDR = 0.15) and FC cutoff 1.2 to the microarray data, 243 genes were found to be up-regulated and 61 probes down-regulated. A volcano plot visualizes the probes that were ruled in for further study [Figure 3a]. Both previously studied genes and novel genes not previously reported were found in the data set [Tables 2a and 3a]. Pathway analysis indicated the enrichment of MAPK signaling and insulin signaling pathways in the cochlear response to 116 dB SPL noise exposure [Table 4a]. Nine MAPK signaling genes, which include *Fos, Jun, Dusp1, Dusp14, Dusp5, Dusp6, Gadd45g, Nr4a1*, and *Srf*, were up-regulated. Subsequent clustering analysis of GO terms relating to molecular function shows enrichment of the GO term "MAPK phosphatase activity" [Table 5a].

To evaluate the changes that occur in 110 dB SPL noise exposure, gene expression immediately following 1 h noise exposure at 110 dB SPL was compared with control (110 vs. control). A total of 155 probes were found to be up-regulated and 221 down-regulated. Regulated probes included both novel and previously studied genes [Tables 2b and 3b]. The greater inclusion of down-regulated probes is apparent on the volcano plot [Figure 3b]. Pathway analysis indicated the enrichment of MAPK signaling and seven other pathways in the cochlear response to 110 dB SPL noise exposure [Table 4b]. Clustering of GO terms relating to molecular function are enriched for (1) MAPK activity, (2) carbohydrate binding and (3) phosphatase activity [Table 5a].

A subset of genes from the enriched data sets was selected for verification by qPCR. Genes selected for qPCR analysis included both novel genes (*Gem, Serpine1, Cyr61, H19, Ntm1, IgJ*) and genes previously identified in microarray studies following noise exposure (*Fos, Egr1, Mt1*). Good correlation was observed between the microarray and qPCR, as both methods showed consistent up-or down-regulation of selected genes [Figure 4a and b]. Out

of the 14 comparisons verified via qPCR, one comparison (using *IgJ* primers) showed a significant difference in FC expression relative to its microarray counterpart. *IgJ* showed 3.9 times greater down-regulation (9.3-FC vs. 2.4-FC) via qPCR than in microarray studies. Since qPCR is a sensitive technique, a greater FC detected with qPCR compared with microarray hybridization for a specific gene is not surprising.

Finally, since we carried out a genome-wide search, we wanted to examine whether the differentially expressed genes in our study were clustered to any specific chromosomes or if any loci had a high density of these genes. We mapped the location of each gene to its chromosome and found that there was no predilection to any one particular chromosome among the genes in our enriched data set [Table 6].

Discussion

Here, we present the first report of gene expression changes in CBA/CaJ mouse cochleae immediately following 1 h of noise exposure. Our study demonstrates that the activation of MAPK signaling is already present at this early time point in the mouse model. Good correlation between microarray and qPCR data validates the results obtained from global gene expression profiling of cochlear RNA after noise exposure as both methods showed consistent up- or down-regulation of all selected genes. The data presented here reveal a novel set of transcripts linked to early responses to noise exposure. This study provides further evidence that despite the activation of common signaling pathways, subtle differences in distinct transcriptomes differentiate the degree of hearing loss after different levels of noise exposure.

Previously identified and novel genes were found in both data sets [Tables 2 and 3]. For 116 dB SPL vs. control, 243 up-regulated and 61 down-regulated genes were identified. Cluster analysis of GO terms relating to molecular function shows enrichment for the GO term "MAPK phosphatase activity" and for pathways signaling transduction, inflammation, cytoprotective and stress response. For 110 dB SPL vs. control, 155 up-regulated and 221 down-regulated genes were identified. Pathway enrichment can be generally divided into categories of signal transduction, inflammation and stress response. We have identified a novel set of transcripts common to both exposures and a unique set of transcripts that distinguish subtle differences in the metabolic process induced by different levels of noise exposure. This observation is consistent with previous reports that compare different intensities of noise exposure.^[2]

Interestingly, both levels of noise exposure appear to activate the MAPK signaling pathway to induce hearing loss, which is in agreement with the previous reports.^[2,26,27] The observations of this study further suggest that, irrespective of the differences in the noise levels MAPK signaling is potentially a common and critical mediator in NIHL. Nevertheless, subtle differences were detected in the expression of some transcriptomes, even though the modulation of many of the signaling proteins in this pathway was similar in both levels of noise exposure. More importantly, the functional implications of the observed quantitative and qualitative differences are consistent with the corresponding differences in the pattern of hearing loss. For example, the noise-induced increases in the expression of

Fos, which has been implicated in neuronal apoptosis and cell death,^[28,29] was relatively lower in animals exposed to 110 dB SPL. This probably limits the hearing loss to a lower level than that observed with 116 dB SPL, in which the hearing threshold was 10–15 dB higher. In addition, the differential expressions of distinct MAPK signaling transcriptomes that facilitate a survival response also correspond with the levels of hearing loss. Particularly, genes that encode calcium channel proteins Cacnalb and Cacnalg and a calcium-independent phospholipase Pla2g6 were down-regulated at lower levels of noise exposure but were not altered at higher levels. Down-regulation of these genes in the cochlea might be part of the endogenous mechanism that is protective against low-level noise exposure. Noise exposure has been reported to favor the entry of calcium into the sensory cells, which in turn facilitates hearing loss.^[30] The down-regulation of these genes could negatively impact this mechanism and contribute to the relatively lower levels of hearing loss observed at lower exposure levels. Taken together, these observations indicate that, despite the activation of common signaling pathways, the differential expression of distinct transcriptomes in MAPK signaling could determine the differences in the pattern of hearing loss after different levels of noise exposure. One important question relevant to the mechanism of NIHL is, "Which cell type(s) within the cochlear duct show differential expression or changes in specific signaling pathways?" Since the global expression study was carried out using whole cochlear tissue, it is not possible to assign cell type specific changes in gene expression or cell specific signaling pathways. One of the goals of a future study will be to investigate cell type specific changes in gene expression with a special focus on hair cells, support cells, ganglion cells and cells of the stria vascularis.

The partial overlap of our data set to previous noise exposure studies from other laboratories and those transcripts showing the same trend (up- or down-regulated) [Table 2] reveals several important points: First, irrespective of the animal model (mouse, chinchillas or guinea pigs), the type of noise exposure (TTS, PTS or impulse noise), or the duration of exposure to noise and/or the postexposure duration before tissue harvest, [13,14,20] our analysis reveals that a set of genes linked to transcriptional regulation, oxidative and inflammatory stress response and cell cycle/differentiation are up-regulated following noise exposure. In contrast, using a similar comparison to differential gene expression profiles in the salicylate ototoxicity of the mouse,^[21] we found only two transcripts that overlapped with our data set. We propose that those genes noted in Table 2 are early-common responders to noise exposure and thus, generic biomarkers for noise exposure. Second, the cross-species consensus noted above suggests that a similar response is likely in humans following noise exposure. Third, noise exposure induces oxidative stress within an hour of exposure. Fourth, partial overlap of transcripts to previously reported studies noted above validates the methodology used to study the consequences to noise exposure at the molecular level.

In summary, different levels of acoustic over-stimulation result in profound changes in numerous common and divergent molecular pathways in as little as 1 h. We know that these pathways play important roles in tissues and organs outside of the ear. Thus, to interfere with the processes leading to different levels of threshold shifts, and therefore, rescue noise-damaged hearing, careful and targeted regulation of these metabolic pathways will be

required. By better understanding the metabolic milieu of the noise-exposed ear, we hope to positively impact the current epidemic of NIHL.

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Figure 1.

Flow diagram for methods used in this study. Total cochlear RNA recovered from three separate groups (116 dB SPL and 110 dB SPL noise exposure, and Control) were used for microarray hybridization and subsequent qPCR analysis. Differentially expressed genes, identified via analysis of variance and Benjamini-Hochberg analyses, were later used for GO studies and pathway analysis



Figure 2.

Noise-induced shift in hearing threshold. In a different experiment, separate groups of mice were noise-exposed using the same equipment and protocol. Auditory brain stem response was tested prior to noise exposure and 11 days postnoise exposure to assess the permanent noise-induced threshold shift. The graph illustrates the shifts in the threshold detected by auditory brainstem responses recorded at 8, 16, and 32 kHz in mice exposed to 110 dB SPL (n = 8) and 116 dB SPL (n = 6) noise (for one hour) in comparison to control (n = 7). Note that the 110 dB exposure produces a smaller threshold shift than the 116 dB exposure. The results are expressed as mean \pm standard deviation



Figure 3.

Volcano plots of microarray data. Analysis of variance was used to identify differentially expressed candidates in each of the comparisons evaluated in this study. This analysis returned a *P* value for the observed difference (i.e., fold-change [FC]) in expression for a given probe. In the following plots, Log2 (FC) on the *X*-axis was plotted against Log10(*P* value) on the *Y*-axis. The red dots indicate probes that were ruled out for further analysis, while the green dots indicate probes that were ruled in for pathway analysis via Benjamini-Hochberg analyses using a false discovery rate (FDR) as indicated: (a) 116 dB SPL versus control-FDR = 0.15; (b) 110 dB SPL versus control-FDR = 0.15



Figure 4.

Comparison between microarray and real time polymerase chain reaction (PCR) results for selected genes. To verify the relative fold-changes (FCs) observed in the microarray analysis, total RNA recovered from each exposure group was used in real-time PCR (qPCR) analyses. Selected genes were evaluated for consistent up- or down-regulation. Relative FCs for each of the three different comparisons are as shown: (a) 116 dB SPL versus control, (b) 110 dB SPL versus control. qPCR assays were run in triplicate

Table 1

Primer sequences of genes selected for qPCR analysis

Symbol	Gene title	PCR Primers	Sequences	Reference ID	Affymetrix ID
Actb	Actin, beta, cytoplasmic	Forward Reverse	TTGCTGACAGGATGCAGAAG AGTCCGCCTAGAAGCACTTG	NM_007393	M12481_3_at
Cyr61	Cysteine rich protein 61	Forward Reverse	TGGACAACCAGTGTGTACAGCA GATGCGGGCAGTTGTAGTTA	BM202770	1438133_a_at
Egr1	Early growth response 1	Forward Reverse	TCACCCACCATGGACAACTA AGCGGCCAGTATAGGTGATG	NM_007913	1417065_at
Fos	FBJ osteosarcoma oncogene	Forward Reverse	GAATGGTGAAGACCGTGTCA TGGGCTGCCAAAATAAACTC	AV026617	1423100_at
Gem	GTP binding protein	Forward Reverse	TAGTGCGGTGTCGAGGAGGTG GACAGGTCATGGCCAGGATTT	U10551	1426063_a_at
H19	H19 fetal liver mRNA	Forward Reverse	AATGGTGCTACCCAGCTCAT GCAGAGTTGGCCATGAAGAT	NM_023123	1448194_a_at
Igi	Immunoglobulin joining chain	Forward Reverse	CCCTTTGAACAACAGGGGAGA AGGTCTCAGGGAACACCATCG	BC006026	1424305_at
Mt1	Metallothionein 1	Forward Reverse	CACCAGATCTCGGAATGGAC TGTCCCCAACAGAGAAGACC	BC027262	1451612_at
Ntn1	Netrin 1	Forward Reverse	TCCAAAGGCAAGCTGAAGAT GGCATTACCCAACAGCAAGT	BI143915	1454974_at
Serpine1	Serine (or cysteine) peptidase inhibitor, clade E, member 1	Forward Reverse	CCGATCCTTTCTCTTTGTGG GGATGCTGGTTGGGAAGGATT	NM_008871	1419149_at

qPCR = Quantitative real-time polymerase chain reaction

Table 2a

Previously reported genes linked to 116 dB SPL noise exposure

Symbol	Gene title	Fold-change	Р	Reference ID
Transcriptiona	l regulators			
Fos ^{<i>a</i>*,<i>b</i>*,<i>d</i>}	FBJ osteosarcoma oncogene	5.6	0.0002	AV026617
Egr1 ^{a*,b*}	Early growth response 1	4.7	0.0001	NM_007913
Nr4a3 ^b	Nuclear receptor subfamily 4, group A, member 3	3.5	0.0005	BE692107
Nr4a1 ^{b*}	Nuclear receptor subfamily 4, group A, member 1	3.3	0.0001	NM_010444
Atf3 ^{b*}	Activating transcription factor 3	3.1	0.0001	BC019946
Jun ^b	Jun oncogene	1.6	0.0164	NM_010591
Zfp36 ^b	Zinc finger protein 36	1.5	0.0075	X14678
Oxidative and	inflammatory stress response			
Dusp1 ^{b*}	Dual specificity phosphatase 1	2.0	0.0002	NM_013642
Dusp5 ^{b*}	Dual specificity phosphatase 5	1.7	0.0015	BB442784
Ptgs2 ^{b*}	Prostaglandin-endoperoxide synthase 2	1.6	0.0014	M94967
Pvr ^b	Poliovirus receptor	1.4	0.0226	BB049138
Cell cycle/differentiation				
Btg2 $a^{*,b}$	B-cell translocation gene 2	1.6	0.0156	NM_007570
Btg2 $a^{*,b}$	B-cell translocation gene 2	1.6	0.0289	NM_007570
Gdf15 ^{b*}	Growth differentiation factor 15	1.2	0.0299	NM_011819

^aPrior NIHL study = 90 min noise exposures (PTS and TTS), 2.0-fold-changes; no FDR utilized (Cho et al. 2004),

^bPrior NIHL study = Impulse noise, 2.0-fold changes (P < 0.008), FDR = 0.25 (Kirkegaard *et al.*, 2006),

^cSalicylate ototoxicity = 2-fold up-regulation; no FDR utilized (Im *et al.*, 2007)

^dPrior NIHL study = 90 min noise exposures (PTS and TTS) (Lomax *et al.* 2001),

* Microarray data confirmed by qPCR

Table 2b

Previously reported genes linked to 110 dB SPL noise exposure

Symbol	Gene title	Fold-change	Р	Reference ID
Transcriptiona	l regulators			
Fos ^{<i>a</i>*,<i>b</i>*,<i>d</i>}	FBJ osteosarcoma oncogene	7.4	0.0001	AV026617
$Egr1^{a^*,b^*}$	Early growth response 1	5.4	0.0001	NM_007913
Nr4a1 ^{b*}	Nuclear receptor subfamily 4, group A, member 1	4.5	0.0000	NM_010444
Nr4a3 ^b	Nuclear receptor subfamily 4, group A, member 3	4.1	0.0002	BE692107
Atf3 ^{b*}	Activating transcription factor 3	4.0	0.0000	BC019946
Cebpb ^b	CCAAT/enhancer binding protein, beta	2.4	0.0001	NM_013154
Maff ^{c*}	V-maf musculoaponeurotic fibrosarcoma oncogene family, protein F	2.1	0.0000	BC022952
Jun ^b	Jun oncogene	2.0	0.0024	NM_010591
Zfp36 ^b	Zinc finger protein 36	1.6	0.0025	X14678
Nab2 ^d	Ngf-A binding protein 2	1.4	0.0255	NM_008668
Oxidative and	inflammatory stress response			
Dusp1 ^{b*}	Dual specificity phosphatase 1	2.2	0.0001	NM_013642
Dusp5 ^{b*}	Dual specificity phosphatase 5	1.8	0.0007	BB442784
Ptgs2 ^{b*}	Prostaglandin-endoperoxide synthase 2	2.0	0.0002	M94967
Pvr ^b	Poliovirus receptor	1.3	0.0043	BB049138
Pvr ^b	Poliovirus receptor	1.2	0.0076	NM_009310
Cell cycle/diffe	erentiation			
Btg2 $a^{*,b}$	B-cell translocation gene 2	1.9	0.0041	NM_007570
Btg2 $a^{*,b}$	B-cell translocation gene 2	1.8	0.0131	NM_007570
Gdf15 ^{b*}	Growth differentiation factor 15	1.4	0.0044	NM_011819

^aPrior NIHL study = 90 min noise exposures (PTS and TTS); 2.0-fold-changes; no FDR utilized (Cho et al. 2004),

^bPrior NIHL study = Impulse noise, 2.0-fold-changes (P < 0.008), FDR = 0.25 (Kirkegaard *et al.* 2006),

^cSalicylate ototoxicity = 2-fold up-regulation, No FDR utilized (Im *et al.* 2007),

^dPrior NIHL study = 90 min noise exposures (PTS and TTS) (Lomax *et al.* 2001),

* Microarray data confirmed by qPCR

Table 3a

Novel genes linked to 116 dB SPL noise exposure

Gene symbol	Gene title	Fold change	Р	Reference ID
Transcriptional regu	ators			
Junb	Jun-B oncogene	2.7	0.0002	NM_008416
Signal transduction				
Gem	GTP binding protein (gene overexpressed in skeletal muscle)	3.1	0.0005	U10551
Extracellular/ECM associated/secreted proteins				
Apold1	Apolipoprotein L domain containing 1	3.5	0.0001	BM123813
Serpine1	Serine (or cysteine) peptidase inhibitor, clade E, member 1	2.9	0.0001	NM_008871
Cyr61 ^{**}	Cysteine rich protein 61	2.3	0.0000	NM_010516
Cyr61 ^{**}	Cysteine rich protein 61	2.1	0.0001	BM202770
Cyr61**	Cysteine rich protein 61	2.0	0.0176	BB533736
Other				
3300001A09Rik	RIKEN cDNA 3300001A09 gene	1.9	0.0226	AK014341
Usp53	Ubiquitin specific peptidase 53	1.9	0.0005	AV320152
Zfp533	Zinc finger protein 533	-1.4	0.0168	BE982894

** Duplicate entries represent different isoforms of the same gene or the same gene identified with a different probe sequence, ECM = Extracellular matrix

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Table 3b

Novel genes linked to 110 dB SPL noise exposure

Gene symbol	Gene title	Fold-change	Р	Reference ID
Transcriptional regulators				
Junb	Jun-B oncogene	3.2	0.0001	NM_008416
Signal transduction				
Gem	GTP binding protein (gene overexpressed in skeletal muscle)	4.7	0.0001	U10551
Extracellular/ECM associated/secreted proteins				
Serpine1	Serine (or cysteine) peptidase inhibitor, clade E, member 1	4.2	0.0000	NM_008871
Apold1	Apolipoprotein L domain containing 1	3.7	0.0000	BM123813
Cyr61 ^{**}	Cysteine rich protein 61	3.1	0.0000	NM_010516
Cyr61 ^{**}	Cysteine rich protein 61	2.8	0.0000	BM202770
Cyr61 ^{**}	Cysteine rich protein 61	2.3	0.0080	BB533736
Cyr61 ^{**}	Cysteine rich protein 61	2.1	0.0117	BB533736
Other				
H19	H19 fetal liver mRNA	-1.7	0.0250	NM_023123

** Duplicate entries represent different isoforms of the same gene or the same gene identified with a different probe sequence, ECM = Extracellular matrix

Table 4a

116 dB SPL noise exposure: Enriched pathways

Term	Count	Percentage	Р
MAPK signaling	9	0.5	2.1E-3
Insulin signaling	6	0.3	7.2E-3

Table 4b

110 dB SPL noise exposure: Enriched pathways

Term	Count	Percentage	Р
MAPK signaling	13	0.6	2.4E-5
Type II diabetes mellitus	4	0.2	1.6E-2
Calcium signaling	7	0.3	1.8E-2
B cell receptor signaling	4	0.2	5.7E-2
Primary immunodeficiency	3	0.1	5.9E-2
Fc epsilon RI signaling	4	0.2	6.1E-2
Wnt signaling	5	0.2	8.0E-2
Fc gamma R-mediated phagocytosis	4	0.2	9.2E-2

Table 5a

116 dB SPL noise exposure: Functional annotation clustering of similar GO terms relating to molecular function

MAPK phosphatase activity*				
EASE score average ^{a} = 3.26	EASE score ^b	Fold enrichment ^c		
GO:0033549~MAPK phosphatase activity	3.27E-04	27.9		
GO:0017017~MAPK tyrosine/serine/threonine phosphatase activity	3.27E-04	27.9		
$GO:0008138 \text{-} protein tyrosine/serine/threonine phosphatase activity}$	1.58E-03	9.8		

^{*a*}EASE score average represents the-Log of the averaged EASE score per cluster, where 1.3 = -Log (P = 0.05),

 b EASE scores are the P values of a modified Fisher's exact test. EASE score <0.05 is considered significant,

^cDegree of GO enrichment relative to background present on Affymetrix Mouse Genome 430 2.0 array,

* Represents the most commonly shared GO term per cluster, GO = Gene ontology

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Table 5b

110 dB SPL noise exposure: Functional annotation clustering of similar GO terms relating to molecular function

MAPK tyrosine/serine/threonine phosphatase activity *				
EASE score average ^{a} = 5.2	EASE score ^b	Fold enrichment ^c		
GO:0017017~MAPK tyrosine/serine/ threonine phosphatase activity	7.11E-07	30.6		
GO:0033549~MAPK phosphatase activity	7.11E-07	30.6		
GO:0008138~protein tyrosine/serine/threonine phosphatase activity	5.91E-04	8.6		
Carbohydrate binding				
EASE score average ^{a} =2.2	EASE score b	Fold enrichment ^C		
GO:0005539~glycosaminoglycan binding	1.08E-03	5.0		
GO:0030247~polysaccharide binding	1.77E-03	4.6		
GO:0001871~pattern binding	2.63E-03	4.3		
GO:0008201~heparin binding	7.97E-03	4.8		
GO:0030246~carbohydrate binding	1.59E-01	1.7		
Phosphatase activity				
EASE score average ^{a} =1.6	EASE score b	Fold enrichment ^C		
GO:0004721~phosphoprotein phosphatase activity	5.71E-03	3.3		
GO:0016791~phosphoric monoester hydrolase activity	2.69E-02	2.4		
GO:0042578~phosphoric ester hydrolase activity	8.79E-02	1.9		

^{*a*}EASE score average represents the-Log of the averaged EASE score, where 1.3 = -Log (P = 0.05),

 b EASE scores are the P values of a modified Fisher's exact test. EASE score <0.05 is considered significant.

 c The degree of GO enrichment relative to background present on Affymetrix Mouse Genome 430 2.0 array,

* Represents the most commonly shared GO term per cluster, GO = Gene ontology

Table 6

Mapping of differentially expressed genes to chromosome

Chromosome	116 versus control	110 versus control
1	20	23
2	27	30
3	26	22
4	19	27
5	11	18
6	23	21
7	16	25
8	10	19
9	14	17
10	16	18
11	21	25
12	15	12
13	13	15
14	6	9
15	14	16
16	8	13
17	10	23
18	8	15
19	8	8
Х	10	10

We mapped the location of every gene from the three enriched data sets to their chromosome. Each number in a column represents the number of genes from a given data set that mapped to a specific chromosome. This data shows that there is no clustering of "NIHL genes" to any one particular chromosome among the genes in our enriched data set.