Auditory Phenotype and Histopathologic Findings of a Mutant Nlrp3 Expression Mouse Model

Yehree Kim1†, Sang-Yeon Lee2†, Min Young Kim1, Kyusun Park1, Jin Hee Han1, Jung Ho Kim3, Bong Jik Kim4* and Byung Yoon Choi1*

1 Department of Otorhinolaryngology, Seoul National University Bundang Hospital, Seongnam, South Korea, 2 Department of Otorhinolaryngology, Seoul National University Hospital, Seoul, South Korea, 3 Department of Pathology, Seoul National University Hospital, Seoul, South Korea, 4 Department of Otorhinolaryngology-Head and Neck Surgery, Chungnam National University, Chungnam National University Sejong Hospital, Sejong, South Korea

Objective: The pathogenesis of hearing loss in autoinflammatory disorders due to activation of the inflammasome remains incompletely understood. Previously no animals expressing mutant Nlrp3 (NOD-, LRR- and pyrin domain-containing protein 3) survived to an age when hearing evaluation was possible due to embryonic lethality. We aimed to establish a novel mouse model that manifests quantifiable hearing loss with other syndromic features due to alteration of Nlrp3 and investigate the audiologic and histopathologic phenotype in the cochlea to clarify how the genetic alterations of NLRP3 could induce autoinflammatory hearing loss.

Methods: To induce inner ear expression of the mutant Nlrp3, Nlrp3D301NneoR mice were bred with Gfi1Cre knock-in mice for conditional mutant Nlrp3 activation in the cochlea and hematopoietic cells. Hearing thresholds were measured. Hematoxylin-eosin sections of the cochlea, brain, kidney, and liver were examined under light microscopy. Immunohistochemical analyses using polyclonal anti-NLRP3 antibodies on cochlear whole-mount preparations and frozen sections were performed.

Results: We, for the first time in the literature, established a mouse model that manifests quantifiable hearing loss due to Nlrp3 alteration. ABR recordings of Nlrp3D301NneoR/+; Gfi1Cre/+ mice, albeit with limited life expectancy, exhibited severe to profound hearing loss at postnatal day 20 (P20). There was overall overexpression of mutant Nlrp3, and mutant Nlrp3 expression was noted in the spiral prominence, the outer sulcus region (Claudius cells and outer sulcus cells), the organ of Corti, the inner sulcus, and the spiral ganglion neurons in the cochlea. Hematoxylin-eosin sections of Nlrp3D301NneoR/+; Gfi1Cre/+ mice cochleae at P12 exhibited a disorganized organ of Corti between the outer hair cells/supporting Deiters’ cells and basilar membrane compared with the normal phenotype mice, leading to a collapsed Nuel’s space. This morphologic feature gradually returned to normal by P15. Varying degrees of inflammation with lymphocytic infiltrations were observed in the brain, kidney, and liver.

Conclusion: We report the first mutant Nlrp3 overexpression mouse model (Nlrp3D301NneoR/+; Gfi1Cre/+)) that shows obvious overexpression of Nlrp3 in the cochlea, a transient developmental lag of the cochlea, and severe to profound hearing loss.
INTRODUCTION

The NLRP3 (NOD-, LRR-, and pyrin domain-containing protein 3) gene encodes the NLRP3 protein, the central component of the NLRP3 inflammasome (1). The NLRP3 inflammasome is an innate immune sensor expressed in immune cells (2, 3), that leads to the secretion of interleukin-1β when activated. Activation of the NLRP3 inflammasome transforms the inactive procaspase-1 to active caspase-1 which can process pro-IL-1β to mature IL-1β (4).

Alterations in the NLRP3 gene are associated with several autoinflammatory disorders affecting several organs, referred to as cryopyrin-associated periodic syndromes (CAPS), which include a spectrum of disorders—chronic infantile neurological cutaneous and articular syndrome (CINCA or NOMID, neonatal onset multisystem inflammatory disease), Muckle-Wells syndrome and familial cold autoinflammatory syndrome (FCAS) (5, 6).

Hearing loss is one of the presenting symptoms of these autoinflammatory disorders. About 76, 86, 33, and 25% of NOMID, NOMID/MWS, MWS, and FCAS subjects exhibit hearing loss (7). Recent reports have suggested that hearing loss could be the sole symptom in a subset of autoinflammatory diseases (DFNA34) (8, 9).

Activation of the NLRP3 inflammasome requires at least two signals—the initial priming signal and the second activation signal (10). However, in patients with pathogenic variants of NLRP3 gene, this process only requires the initial priming signal to induce IL-1β secretion (11). Anakinra, a non-glycosylated recombinant version of the endogenous human IL-1 receptor antagonist, was reported to control systemic inflammation (12). Intriguingly, it also showed partial effectiveness in the treatment of autoinflammatory hearing loss (9).

Cochlear implantation in patients with this disease entity demonstrated successful auditory rehabilitation outcomes (13). Cochlear implantation could work as a rescue measure in cases of advanced, profound NLRP3-related hearing loss, and an anti-IL-1β agent could be beneficial for early diagnosed, mild hearing loss. However, it has not been clearly elucidated how hearing loss develops in this disease entity; therefore, prevention or timely intervention remains a challenging issue in the management of hearing loss as a presenting symptom of autoinflammatory disorders.

A prior study identified Nlrp3 expression in normal mouse cochlear macrophage-like cells expressing Cx3cr1 (8). The authors found that the NLRP3 inflammasome could be activated with the priming and the activating signals, thus indicating that macrophage/monocyte-like cells in the cochlea can cause hearing loss through innate immune response.

To study the pathogenesis of hearing loss in autoinflammatory disorders due to activation of the inflammasome from developmental and immunological perspectives, a mouse model recapitulating human autoinflammatory disorders with quantifiable hearing loss is mandatory. However, in a previous model, severe inflammation due to Nlrp3 mutation resulted in an extremely short life expectancy, which made an objective hearing assessment impossible.

We generated a double knock-in mouse model by breeding Nlrp3<sup>D301NneoR</sup> knock-in mice with Gfi1<sup>Cre</sup> (growth factor independent I transcriptional repressor) knock-in mice, a known tool for inducing inner ear hair cell-specific expression of a target gene (14–16) thereby generating the first-ever mouse model in the literature manifesting with both hearing loss and overexpression of Nlrp3. In this study, first, we conducted detailed audiologic and histopathologic phenotyping of gain of function of mutant Nlrp3 in the cochlea. By doing this, we aimed to elucidate how alterations of NLRP3 in the cochlea could play a role in the development of autoinflammatory hearing loss.

MATERIALS AND METHODS

Animals and Housing

To generate the Nlrp3 gain-of-function model, we bred congenital-onset, multisystemic inflammatory disease mice (Nlrp3<sup>D301NneoR</sup>, obtained from The Jackson Laboratory, Jax #017971) with heterozygous Gfi1<sup>Cre</sup> knock-in mice (15) to generate mice mutant (Nlrp3<sup>D301NneoR</sup>+/+; Gfi1<sup>Cre</sup>/+) and normal phenotype littermate controls (Nlrp3<sup>D301NneoR</sup>+/+; Gfi1<sup>Cre</sup>/+). An F1 hybrid mouse was generated on a C57BL/6 background and Gfi1<sup>Cre</sup> knock-in mice on 129S6 and C57BL/6 background (17). All animals were kept on a 12-h light-dark cycle in a specific-pathogen-free facility with controlled temperature and humidity and had free access to food and water. All experimental procedures were conducted according to protocols approved by the Institutional Animal Care and Use Committee of Seoul National University Bundang Hospital (Seongnam, Republic of Korea, IACUC No.: BA-2004-294-038-09).

Conditional Expression of Mutant Nlrp3 in the Mouse Inner Ear

To induce inner ear expression of the mutant Nlrp3, Nlrp3<sup>D301NneoR</sup> mice bearing a loxP-flanked neomycin-resistant
cassette (reverse orientation) in intron 2 and a point mutation in exon 3 of Nlrp3 were bred with Gfi1Cre knock-in mice, generating Nlrp3D301NeoR /+; Gfi1Cre/+ mice for conditional mutant Nlrp3 activation only in the cochlea and hematopoietic cells.

Genotyping was performed by polymerase chain reaction (PCR) on genomic DNA extracted from ear punches. The primers used for Gfi1Cre genotyping were as follows: Gfi1 wild-type forward (5′-GGG ATA ACG GAC CAG TTG-3′), Gfi1 wild-type reverse (5′-CCG AGG GGC GTT AGG ATA-3′), and Gfi1Cre reverse (5′-GCC CAA ATG TTG CTP AGT-3′). The primers used for Nlrp3 genotyping were as follows: Nlrp3 wild-type forward (5′-CAC CCT GCA TTT TGT TG-3′), Nlrp3 mutant forward (5′-GCT ATT TGT CAC GTC C-3′), and common reverse (5′-CGT GTA GCG ACT GTT GAG GT-3′).

Auditory Brainstem Responses

The hearing threshold of 10 Nlrp3D301NeoR /+; Gfi1Cre/+ mice (seven males and three females), six Nlrp3D301NeoR /+; Gfi1Cre/+ mice (four males and two females), four Nlrp3+/++; Gfi1Cre/+ mice (two males and two females) and four Nlrp3+/++; Gfi1Cre/+ mice (two males and two females) were measured at postnatal day 20 (P20). The hearing threshold of the right ear of each animal was used for analysis. For auditory brainstem response (ABR) testing, mice were anesthetized with an intraperitoneal injection of a mixture of zoalazepam/tiletamine (50 mg/kg, Zoletil 50; Virbac, Carros, France) and 10 mg/kg xylazine (Rompun; Bayer Health Care, Leverkusen, Germany). Reference electrodes were attached posterior to the right pinna, a non-inverting electrode was attached at the vertex of the skull, and a ground electrode was attached to the contralateral ear (17). Hearing thresholds were determined at 4, 8, 16, and 32 kHz using a SmartEP device (Intelligent Hearing Systems, Miami, FL, USA). Acoustic stimuli were presented directly to the entrance of the ear canal. Tone burst sound stimuli were presented from 100 dB SPL and decreased 10 dB decrements until the lowest level at which a distinct ABR wave pattern could be recognized by two of the investigators. The ABR threshold was determined as the lowest recognizable ABR response. For mice with no ABR responses, the hearing threshold was calculated as 110 dB SPL. All recordings were performed in a soundproof box, and ABR was tested at postnatal day 20 (P20).

Acquisition of Magnetic Resonance Images

To check for cochlear inflammation, pre- and post-contrast enhanced magnetic resonance imaging (MRI) was performed (n = 2, one Nlrp3D301NeoR /+; Gfi1Cre/+ and one Nlrp3D301NeoR /+; Gfi1Cre/+; both male) at P15. All MR scans were performed on a MRS’ DRYMAG 7 Tesla (T) MRI system (MR solutions, Guilford, UK) using gradients with a maximum gradient strength of 420 mT/m. Mice were anesthetized as described in the ABR section and body temperature was maintained at 37°C by circulating warm air through the magnet bore. T1-weighted images were acquired first. The mice were scanned using a twodimensional (2D) spoiled gradient echo sequence (acquisition matrix = 256 × 256; field of view = 20 mm × 20 mm; slice thickness = 3 mm; flip angle = 90°).

Immediately after T1 measurement, the mouse was removed from the scanner, injected with a single bolus of gadolinium meglumine (Dotarem; Guerbet, Villepinte, France) at a dose of 0.2 mmol/kg through the tail vein, and then re-positioned in the scanner. Proper delivery of contrast agent was confirmed first by scanning the abdomen and checking for contrast material in the mouse bladder. Thirty minutes after the injection of contrast agent, contrast enhanced images were acquired using the same acquisition matrix as the precontrast images.

Cochlear Exraction and Preparation of Histologic Sections

To examine the presence of any cochlear structural abnormalities, hematoxylin-eosin stained was performed at P12, P15, and P20 (n = 4 cochleae of Nlrp3D301NeoR /+; Gfi1Cre/+ mice, and n = 4 cochleae of Nlrp3D301NeoR /+; Gfi1Cre/+ for each timepoint). Mice were anesthetized as described before in the ABR section and decapitated. Both cochleae were extracted and fixed in ice-cold 4% paraformaldehyde overnight. The fixed cochleae were washed, dehydrated, and embedded in paraffin. Serial sections were made at 5-µm thickness, attached to the slide glass, and air-dried. The tissue sections were deparaffinized, hydrated, stained with HE, dehydrated, cleared, and sealed with a cover glass. The histologic characteristics were examined under a light microscope.

Immunohistochemistry of the Cochlear Tissue

Immunohistochemistry was performed on Nlrp3D301NeoR /+; Gfi1Cre/+ and Nlrp3D301NeoR /+; Gfi1Cre/+ mice at P17 (n = 4 cochleae for each type). Cochlear turns were carefully excised, and whole-mount preparations were made and incubated in blocking/permeabilizing buffer (phosphate-buffered saline [PBS] with 5% goat serum and 0.25% Triton X-100). Cochlear frozen sections were also prepared and incubated in blocking/permeabilizing buffer. The preparations were incubated overnight at 4°C with 1:100 rabbit anti-NLRP3 (PA5-88709, Abcam, Cambridge, UK) secondary antibodies diluted in blocking/permeabilizing buffer. After 3 washes, the cochlear turns were reacted with fluorescence-labeled 1:400 goat anti-rabbit (ab150077, Abcam, Cambridge, UK) secondary antibodies diluted in blocking/permeabilizing buffer for 2 h at room temperature. After 3 washes with PBS, the cochlear turns were reacted with 1:100 rhodamine phalloidin (R415, Invitrogen, Waltham, MA, USA) diluted in blocking/permeabilizing buffer for 1 h at room temperature. The samples were then rinsed 3 times with PBS. High-resolution images were obtained using a confocal laser scanning microscope (LSM710, Zeiss). Each image stack was reduced to a two-dimensional maximum intensity Z-projection image.

Prior to the immunohistochemistry procedures on cochlear tissues, the anti-NLRP3 antibody was validated on human embryonic kidney (HEK) 293 cells transfected with
NLRP3 (Myc-DDK-tagged) cDNA (catalog no. RC220952) (Supplementary Figure 1).

Hair Cell Count
The number of inner and outer hair cells were counted from the confocal images of the whole-mount preparations described above (Immunohistochemistry of the cochlear tissue). Serial confocal images of hair cells labeled with phalloidin were obtained from each cochlea using laser scanning confocal microscope (LSM710, Zeiss). The entire length of the organ of Corti was reconstructed by overlapping the common cells at the edges of the individual images. The reconstructed image was then divided into segments, each spanning 1% of the total length. The number of IHCs and OHCs in each segment vs. the relative distance from the apex were plotted (18).

Histologic Examination of Other Organs
Histologic evaluation with HE staining was performed on two Nlrp3 D301NneoR /−; Gfi1 Cre/+ mice and one Nlrp3 D301NneoR /+; Gfi1 Cre/+ mouse at P20 to check for histopathologic signs of systemic inflammation in other organs. HE-stained tissue slides of the brain, eyes, liver, spleen, stomach, kidney and legs were prepared and the findings were assessed by an experienced pathologist (JHK) with more than 15 years of diagnostic experience. The type of immune cells infiltrating each tissue was determined by the microscopic morphology.

Data Analysis and Statistical Analysis
Data analysis was carried out using SPSS version 25.0 (IBM Corp., Armonk, NY, USA). For comparisons between two samples, the significance of data was assessed by the Student t-test when samples showed a normal distribution or by the Mann-Whitney test when samples did not pass the normality test. Comparisons of more than two samples were conducted using one-way analysis of variance with the Tukey post-hoc test when data passed the normality test or the Kruskal-Wallis test. The difference between the results of the Nlrp3 D301NneoR /+; Gfi1 Cre/+ and the Nlrp3 D301NneoR /−; Gfi1 Cre/+ mice means that the hearing loss observed in the Nlrp3 D301NneoR /+; Gfi1 Cre/+ mice was entirely due to the p.D301N variant and not due to Gfi1 haploinsufficiency.

RESULTS

Auditory Brainstem Responses Reveal Profound Deafness in Nlrp3 D301NneoR /+; Gfi1 Cre/+ Mice
The hearing phenotype was assessed at P20 by ABR recordings, which revealed that Nlrp3 D301NneoR /+; Gfi1 Cre/+ mice (n = 10 mice) exhibited severe to profound hearing loss. For comparison, the ABR thresholds of Nlrp3 D301NneoR /+; Gfi1 Cre/+ mice (n = 4), Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (n = 6) and Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (n = 4) were recorded at P20.

When compared to the ABR thresholds for Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (four frequency average 37.8 ± 13.9 dB), Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (37.1 ± 7.5) and Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (39.4 ± 13.8 dB), the ABR thresholds of the Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (96.4 ± 20.4 dB) were significantly higher across the four tested frequencies (Figure 1, P < 0.05, Kruskal-Wallis test with the Dunn post-hoc test). The difference between the results of the Nlrp3 D301NneoR /+; Gfi1 Cre/+ and the Nlrp3 D301NneoR /−; Gfi1 Cre/+ mice showed similar signal intensity of the cochlea (Figures 2A,C). The same images were acquired after injection of contrast material into the tail vein which showed enhancement of the fluid compartment of the cochlea of the Nlrp3 D301NneoR /+; Gfi1 Cre/+ (Figure 2D). As for the Nlrp3 D301NneoR /+; Gfi1 Cre/+ mouse, there was no enhancement of the cochlea (Figure 2B).

Of note, middle ear inflammation was also visible on the pre- and post-contrast enhanced images of the Nlrp3 D301NneoR /+; Gfi1 Cre/+ mouse (Figures 2C,D).

Localisation of Nlrp3 in the Inner Ear From Nlrp3 D301NneoR /+; Gfi1 Cre/+ and Nlrp3 D301NneoR /+; Gfi1 Cre/+ Mice
To specify Nlrp3 expression in the cochlear tissue, we performed immunohistochemical staining using anti-NLRP3 antibody. The expression of Nlrp3 protein was evaluated in the cochleae of Nlrp3 D301NneoR /+; Gfi1 Cre/+ and Nlrp3 D301NneoR /+; Gfi1 Cre/+ at P15 (Figures 3, 4). In the cochlea of Nlrp3 D301NneoR /+; Gfi1 Cre/+ mouse, which was expected to show
compared in the spiral ganglion neurons (Figure 4B). Images taken 30 min after gadolinium injection, show enhancement of the cochlea in the Nlrp3 D301NneoR /+; Gfi1Cre/+ mouse (D) but not in the cochlea of the Nlrp3 D301NneoR /+; Gfi1Cre/+ mouse (B). The Nlrp3 D301NneoR /+; Gfi1Cre/+ mouse also has inflammation in the middle ear (white arrows) where in the healthy condition should contain air and appear black as can be seen in the middle ear of the Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (asterisks).

The number of inner and outer hair cells were quantified and compared in Nlrp3 D301NneoR /+; Gfi1Cre/+ (A) and the Nlrp3 D301NneoR /+; Gfi1Cre/+ mouse (C). In the organ of Corti (Figures 3A,C), and the spiral ganglion neurons (Figures 3A,B). In the cochlea of Nlrp3 D301NneoR /+; Gfi1Cre/+ mouse, there appeared an overall overexpression of mutant Nlrp3, and the immunostaining was noted in the spiral prominence, the outer sulcus region (Claudius cells and outer sulcus cells) (Figures 3E,H), the organ of Corti, the inner sulcus (Figures 3E,G), and the spiral ganglion neurons (Figures 3E,H). Similar immunostaining patterns of Nlrp3 were seen throughout the whole cochlea along the tonotopic axis (Figures 3I,J). Noticeably, the Nlrp3 expression pattern was shown to differ the most in the outer sulcus region and the inner sulcus between the two types of mice.

Meanwhile, immunohistochemical staining of the whole mount preparations revealed similar but rather restricted expression pattern compared with those observed in the frozen section. The overexpression of mutant Nlrp3 compared to the pattern of wild-type Nlrp3 expression was most prominent at the Hensen’s cells and the outer sulcus region (Figure 4A), as well as the spiral ganglion neurons (Figure 4B).

Hair Cell Count

The number of inner and outer hair cells were quantified and compared in Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (n = 6 cochleae) and Nlrp3 D301NneoR /+; Gfi1Cre/+ mice (n = 6 cochleae) at P15. Neither HIC nor OHC number showed significant difference in these two groups (Supplementary Figure 2). All inner and outer hair cells looked morphologically intact in Nlrp3 D301NneoR /+; Gfi1Cre/+ mice while the ABR results showed profound hearing loss in this group.

Cochlear Histopathology in Nlrp3 D301NneoR /+; Gfi1Cre/+ Mice

A close evaluation of the organ of Corti demonstrated a difference in histopathology between Nlrp3 D301NneoR /+; Gfi1Cre/+ and Nlrp3 D301NneoR /+; Gfi1Cre/+; Gfi1Cre/+ (normal phenotype) mice at P12 (Figure 5). At the middle and apical turns of the cochlea, hair and supporting cells were present in mice of both genotypes. However, Nlrp3 D301NneoR /+; Gfi1Cre/+ mice consistently exhibited a severely collapsed or disorganized organ of Corti between outer hair cells/supporting dieter’s cell and the basilar membrane compared with Nlrp3 D301NneoR /+; Gfi1Cre/+ mice, leading to an apparently collapsed Nuel’s space (space between outer pillar cells and outer hair cells/dieter’s cell). Additionally, nuclei of the greater epithelial ridge and other supporting cells close to the spiral ligament in Nlrp3 D301NneoR /+; Gfi1Cre/+ mice tended to be more condensed than in Nlrp3 D301NneoR /+; Gfi1Cre/+ mice. Meanwhile, no meaningful differences were found in histopathology between Nlrp3+/+; Gfi1Cre/+ and...
**FIGURE 3** | Inner ear section immunohistochemistry from P15 Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> and Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> mice stained with an anti-NLRP3 antibody. Nlrp3 was more widely overexpressed in the Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> mouse (E–H) than in the Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> mouse (A–D). In both types of mice, Nlrp3 was expressed in the spiral prominence (D,H), the organ of Corti (C,G), and the spiral ganglion neurons (B,F). A difference in the expression of Nlrp3 was noted in the inner sulcus cells (arrows), Hensen’s cells (asterisks), and the outer sulcus region (arrowheads). Similar expression pattern of Nlrp3 was observed in two other cochlear turns of Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> (I) and Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> mice (J). n = 6 cochleae for Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> Nlrp3 in green, DAPI in blue, the scale bars represent 100 µm for (A,E,I,J) and 20 µm for (B–D,F–H). P15, postnatal day 15.

*Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> mice, implying that the collapsed morphology of the organ of Corti was also not attributable to Gfi1 haploinsufficiency (Supplementary Figure 3). Interestingly, the sections of *Nlrp3 <sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> mice tended to gradually restore normal morphology of the organ of Corti at P15, as compared to P12, ultimately revealing normal cochlear morphology in most *Nlrp3<sup>D301N</sup> /+; Gfi1<sup>Cre/+</sup> mice at P20 (Figure 6A). The proportion of collapsed organ of Corti at P12 was 100%, which gradually decreased to 21.1% at the apex, 16.7% at the middle and 0% at the base at P20 (Figure 6B). Together, the data indicate that Nlrp3 gain-of-function might predispose mice to a developmental delay of the organ of Corti, especially at the early postnatal stage.

**Systemic Inflammation in the Other Organs of Nlrp3<sup>D301NneoR</sup> /+; Gfi1<sup>Cre/+</sup> Mice**

Various degrees of inflammation were observed on H&E-stained tissue slides of the brain, kidney, and liver (Figure 7). In detail, mild to moderate perivascular lymphocytic infiltration was observed in the brain parenchyma (representative image of the thalamus is shown in Figure 7), and increased inflammatory infiltrates were observed in the renal parenchyma. There was also moderate lymphocytic infiltration in the portal and lobular areas of the hepatic parenchyma. In the spleen, the lymphoid structure seems to be effaced, but definite inflammation was not seen. Definite inflammation was not observed in sections of the eye, stomach, arm, and leg.
FIGURE 4 | Whole-mount immunofluorescence of a P15 Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} mouse cochlea and a Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} mouse cochlea. Nlrp3 was overexpressed in the Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} mouse (bottom panels of (A,B)) compared to the Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} mouse (top panels of (A,B)) in Hensen’s cells and the outer sulcus region (A) and the spiral ganglion neurons (B). (n = 6 cochleae for Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} and n = 6 cochleae for Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}}) Nlrp3 in green, phalloidin in red, DAPI in blue, the scale bars represent 10 µm. P15, postnatal day 15.

DISCUSSION

We, for the first time, report the detailed quantifiable auditory and cochlear histopathologic phenotype of Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} mice. In this animal model, Nlrp3 expression was driven by the Gfi1 promoter, which might be different from the endogenous Nlrp3 expression. Nonetheless, the Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} mouse is the first mouse model to show overexpression of Nlrp3 in the cochlea and potentially associated hearing loss, which could also provide a valuable tool to investigate the link between mutant Nlrp3 overexpression and hearing loss, further revealing the underlying mechanism of immune-related pathophysiology of autoinflammatory hearing loss.

Originally, we intended to generate inner ear-specific expression of Nlrp3 (D301N) by crossing Nlrp3\textsuperscript{D301NneoR/++; Gfi1\textsuperscript{Cre/+}} onto Gfi1-Cre knock-in mice, which was reported as a useful tool for inducing inner ear hair cell-specific expression of a target gene (15). However, Gfi1 encodes a zinc-finger transcription factor that is also important for the development and maintenance of hematopoiesis, which could interfere with maturation of the immune system in the neonatal period and early infancy.
in mice. Accordingly, *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice in our study grew poorly (Supplementary Figure 4) and did not survive longer than 3 weeks (mean survival: 17.5 ± 2.7 days, *n* = 24), which might have been partly due to the intrinsic effect of systemic *Nlrp3* alterations corresponding to the phenotypes observed in human *NLRP3*-related autoinflammatory disorders, such as CINCA, which presents with fever, rash, joint symptoms, and central nervous system symptoms (19). However, its effects might be potentiated by systemic effects on several organs due to the abnormal development of the immune system, as was observed in comparisons of HE staining of several organs between *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* and *Nlrp3*^D301NneoR/+; *Gfi1^+/+* mice.

The widespread inflammation and very short life expectancy observed in the animal model may limit the impact of this study, which only yielded insights into the early changes of *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice in terms of systemic effects and developmental dysregulation of the cochlea and immune system. For instance, the number and morphology of hair cells of *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice were well-preserved not differing from the hair cells of the *Nlrp3*^D301NneoR/+; *Gfi1^+/+* mice. It is possible that the mice in this study did not survive long enough for hair cell damage from prolonged cochlear inflammation to manifest. Future studies using a different Cre mouse that generates milder phenotype and longer survival could show different outcomes. Based on the finding that only around 30% of patients with autoinflammatory inner ear disease respond to anakinra therapy (20) and the hearing in others deteriorate to the level that requires cochlear implantation, the damage caused by the inflammation on the cochlea can be progressive, irreversible and could lead to eventual hair cell damage.

The hearing thresholds of *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice were clearly worse than those of normal controls (Figure 1). A previous study regarding *Gfi1^Cre/+* mice demonstrated early-onset mild progressive hearing loss, which might serve as a source of bias when interpreting the ABR results (14). However, the difference in hearing thresholds between *Gfi1^Cre/+* mice and wild-type (*Gfi1^+/+*) mice in that study was limited to 32 kHz until 2 months of life, meaning that intrinsic hearing deficit due to haploinsufficiency of *Gfi1* did not have a significant impact within <1 month of life. Accordingly, there was no difference in hearing thresholds between *Nlrp3*^+/+*; *Gfi1^Cre/+* mice and *Nlrp3*^D301NneoR/+; *Gfi1^+/+* mice (P > 0.05 at all frequencies by the Mann-Whitney test), although the mean hearing threshold of *Nlrp3*^+/+*; *Gfi1^Cre/+* mice at 32 kHz appeared slightly higher than those of *Nlrp3*^D301NneoR/+; *Gfi1^+/+* mice. On the contrary, *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice showed significantly worse hearing than mice expressing normal *Nlrp3*, irrespective of the loss of one copy of the *Gfi1* gene (Figure 1).

Middle ear inflammation observed in the MR images of the *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice could also have contributed to the elevated hearing thresholds. Indeed, prior studies of ABR threshold measurement in mouse models of otitis media have quantified at most 15 to 30 dB threshold shift depending on the tested frequency (21, 22). However, six out of the 10 *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+* mice in this study did not show any ABR responses in all four tested frequencies even to the strongest stimuli (100 dB SPL). Thus, it is conceivable that a substantial portion of the elevation of hearing thresholds was due to the inner ear dysfunction, not the middle ear inflammation.

Cochlear inflammation was evident on the contrast-enhanced MR images of the *Nlrp3*^D301NneoR/+; *Gfi1^Cre/+*
Kim et al. Mutant Nlrp3 Expression Mouse Model

**FIGURE 6** | Chronological evaluation of cochlear morphology in Nlrp3<sup>D301NneoR<sup>/+</sup>; Gfi1<sup>Cre<sup>/+</sup> mice.

(A) Representative sections from the cochlear turns obtained at P12, P15, and P20, respectively, are depicted (normal cochlear morphology vs. collapsed morphology of organ of Corti).

(B) Proportion of collapsed organs of Corti at each time point, depending on cochlear turns. The scale bars represent 500 mm (P12) and 1,000 mm (P15 and P20). P, postnatal day.
FIGURE 7 | Generalized inflammation observed in several organs of Nlrp3 D301NneoR+/+; Gfi1Cre+ mice compared to Nlrp3 D301NneoR+/+; Gfi1+/+ mice. Representative image of inflammation at P20 in (A) brain (thalamus), (B) kidney, and (C) liver sections of Nlrp3 D301NneoR+/+; Gfi1Cre+ mice (left panels) compared to normal findings in Nlrp3 D301NneoR+/+; Gfi1+/+ mice (right panels). Diffuse lymphocytic infiltration can be seen throughout the whole section. Insets show enlarged views of the dotted areas. Black arrows denote examples of areas with lymphocytic infiltration. P20, postnatal day 20. The scale bars represent 100 µm.

mouse. MRI studies of patients with CAPS or non-syndromic autoinflammatory inner ear disease have also identified cochlear enhancement (7–9). Intravenously injected contrast material can reach the inner ear via small vessels that supply the labyrinth and cochlea (23). Capillary networks present in the stria vascularis and spiral ligament form the blood-labyrinth barrier, composed of endothelial cells with tight-junctions surrounded by pericytes and resident macrophages (24). During inflammatory process of the cochlea, elevated levels of cytokines can increase capillary permeability thereby compromising the integrity of the blood-labyrinth barrier (25). The contrast material can enter the perilymphatic and endolymphatic spaces more freely than in the non-inflammatory state.

Immunohistochemical analyses using the anti-human NLRP3 polyclonal antibody revealed the widespread presence of Nlrp3, not limited to expression in the spiral ganglion, which has been reported in the literature (26). Rather, in the cochlea of Nlrp3 D301NneoR+/+; Gfi1Cre+/+ mice, Nlrp3 was expressed strongly in the spiral prominence, the organ of Corti, and the spiral ganglion neurons. Nlrp3 D301NneoR+/+; Gfi1Cre+/+ mice carry one copy of a wild-type Nlrp3 allele that would result in similar expression of Nlrp3 as seen in Nlrp3 D301NneoR+/+; Gfi1Cre/− mice. The other floxed Nlrp3 allele would lead to the overexpression of Nlrp3 in Cre-expressing cells. Although Gfi1 is known to be expressed in hair cells in the late embryonic and postnatal inner ear (27), a previous validation of a Gfi1Cre/+ mouse model found that Cre-mediated recombination was not entirely hair cell-specific, but was also present in CD45+ monocytes/macrophages (14). A cell type-specific RNA-sequencing study of gene expression during mouse inner ear development also identified Gfi1 expression in...
the surrounding cells of the cochlea at P4 (28). Thus, in the current study, overexpression of Nlrp3 was observed in cells in which Gfi1 was once expressed during the developmental period—that is, in hair cells and the regions of the cochlea where CD45+ cells can be found (in the organ of Corti, the basilar membrane, the spiral ligament, spiral limbus, and the neural region) (29).

The immunoreactivity to Nlrp3 in the cochlear sections differed the most in the inner sulcus and the outer sulcus region between the wildtype and mutant mice, displaying the significant staining exclusively from the mutant mice. The greatest difference in immunoreactivity to Nlrp3 in the cochlear sections between the wildtype and mutant mice was at the region of the inner sulcus and the outer sulcus, with significant positive staining occurring exclusively in the mutant mice. In the spiral prominence and the spiral ganglion neurons, both types of mice showed some degree of immunoreactivity, but it was stronger in the Nlrp3 D301NneoR+/+; Gfi1Cre/+ mice. The aforementioned areas in which we observed differences in immunoreactivity, we could attribute the findings to the ramifications of the gain-of-function variant of Nlrp3. Contrastingly, in the hair cells, the degree of Nlrp3 immunostaining was similar in both types of mice, precluding differentiation of the true immunostaining from staining artifacts. Absence of immunostaining of Nlrp3 from hair cells in the whole mount preparations—unlike in the frozen section—further complicated interpretation of immunostaining in the hair cells. This issue awaits further clarification through validation of the findings on a knockout mouse, which will be done in forthcoming studies.

Il-1β, a downstream product of Nlrp3 inflammasome activation (8), is known to form a positive feedback loop that potentially primes nearby cells in a paracrine manner (30, 31). In the inner sulcus and the outer sulcus region, Gfi1 expression was not anticipated, but overexpression of Nlrp3 was nonetheless evident. In these sites, the induced Il-1β from neighboring cells expressing mutant Nlrp3 could have enhanced the priming step to increase endogenous Nlrp3 expression. Future studies including co-localization of Nlrp3 with Il-1β and treatment with Il-1β agonists would provide further insights.

Intriguingly, as was seen in Figure 6, the morphology of the organ of Corti in Nlrp3 D301NneoR+/+; Gfi1Cre/+ mice gradually normalized from P12 (significant difference between Nlrp3 D301NneoR+/+; Gfi1Cre/+ and Nlrp3 D301NneoR+/+; Gfi1+/+ mice) to P20, which might suggest that the effect of Nlrp3 D301NneoR+/+ on the development of inner ear is transient in the early postnatal period and further indicate that the elevation of the hearing threshold might not be solely due to the anatomical alterations. In further research, we need to generate a new mouse model by breeding Cre mice of specific immune cell population with Nlrp3 knock-in mice, which could express Nlrp3, thereby coming one step closer to endogenous Nlrp3 expression. By doing so, we could elucidate the specific immune cell population associated with the development of autoinflammatory hearing loss in an animal model with a longer lifespan, which could enable a more detailed characterization of hearing loss and immune system dysregulation.

CONCLUSION

Nlrp3 D301NneoR /+; Gfi1Cre/+ mice, an animal model of Nlrp3 expression driven by the Gfi1 promoter, is the first mouse model to show quantifiable hearing loss and overexpression of Nlrp3 in the cochlea. This overexpression is potentially associated with hearing loss, and this model could also provide a valuable tool to investigate the link between mutant Nlrp3 overexpression and hearing loss, further revealing the underlying mechanism of inflammasome activation-mediated hearing loss.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

The animal study was reviewed and approved by Institutional Animal Care and Use Committee of Seoul National University Bundang Hospital.

AUTHOR CONTRIBUTIONS

BK and BC contributed to the conception and design of the study. YK, MK, and JH organized the data and conducted the experiments. YK, S-YL, JK, BK, and BC performed the data analysis. YK, S-YL, BK, and BC wrote the first draft of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

FUNDING

This study was supported by the Basic Science Research Program through the NRF funded by the Ministry of Education (Grants 2021R1A2C2092038 and 2018R1A2B2001054 to BC), the Korea Government (MSIT) (No. 2021R1C1C1007980 to BK), also by SNUBH Intramural Research Funds (16-2020-0005 and 14-2021-0003 to BC), and the Institute of Information Communication Technology Planning and Evaluation (IITP) grant funded by the Korea Government (MSIT) [No. 2020-0-01441, Artificial Intelligence Convergence Research Center (Chungnam National University) to BK].

ACKNOWLEDGMENTS

Gfi1Cre knock-in mice and Nlrp3 D301NneoR knock-in mice were kindly provided for this study by Dr. Jinwoong Bok at the Department of Anatomy, Yonsei University and by Dr. Je-Wook Yu at the Department of Microbiology and Immunology, Yonsei University, respectively.
SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fneur.2022.890256/full#supplementary-material

References

1. Agostini L, Martinon F, Burns K, McDermott MF, Hawkins PN, Tschopp JF. Nlrp3 forms an IL-1β-processing inflammasome with increased activity in Muckle-Wells autoinflammatory disorder. Immunity. (2004) 20:319–23. doi:10.1016/S1074-7613(04)00046-9
2. Sutterwala FS, Ogura Y, Szczepanik M, L INTERNATIONAL IMMUNOLOGY

Supplementary Figure 1 | Validation of the anti-NLRP3 antibody on human embryonic kidney (HEK) 293 cells transfected with NLRP3 (MyoD-DKX-tagged) cDNA. The anti-NLRP3 antibody showed specific binding to the HEK 293 cells transfected with NLRP3 (red color in the bottom panel) and not to the mock transfection control (top panel).

Supplementary Figure 2 | Hair cell count between Nlrp33010WF/+; Gfi1Cre/+ mice and Nlrp33010WF/+; Gfi1Cre/+ mice. Number of inner (A) and outer (B) hair cells in segments spanning 1% of the whole cochlear length in P15 Nlrp33010WF/+; Gfi1Cre/+ mice (n = 6 cochleae) and Nlrp33010WF/+; Gfi1Cre/+ mice (n = 6 cochleae) were not different.

Supplementary Figure 2 | Hair cell count between Nlrp33010WF/+; Gfi1Cre/+ mice and Nlrp33010WF/+; Gfi1Cre/+ mice. Number of inner (A) and outer (B) hair cells in segments spanning 1% of the whole cochlear length in P15 Nlrp33010WF/+; Gfi1Cre/+ mice at P15. Sections of the cochlea of Nlrp33010WF/+; Gfi1Cre/+ mice did not show collapsed morphology of the organ of Corti, unlike the Nlrp33010WF/+; Gfi1Cre/+ mice. P15, postnatal day 15.

Supplementary Figure 4 | Differences in the morphology of Nlrp33010WF/+; Gfi1Cre/+ mice compared to Nlrp33010WF/+; Gfi1Cre/+ mice. At P11, two Nlrp33010WF/+; Gfi1Cre/+ mice were smaller (2.8 g and 3.1 g vs. 6.8 g) and showed more extensive areas of skin inflammation with hair loss than the Nlrp33010WF/+; Gfi1Cre/+ mouse (left). P11, postnatal day 11.
29. Dong Y, Zhang C, Frye M, Yang W, Ding D, Sharma A, et al. Differential fates of tissue macrophages in the cochlea during postnatal development. *Hear Res.* (2018) 365:110–26. doi: 10.1016/j.heares.2018.05.010

30. Gritsenko A, Green JP, Brough D, Lopez-Castejon G. Mechanisms of Nlrp3 priming in inflamming and age related diseases. *Cytokine Growth Factor Rev.* (2020) 55:15–25. doi: 10.1016/j.cytogfr.2020.08.003

31. Hiscott J, Marois J, Garoufalis J, D’addario M, Roulston A, Kwan I, et al. Characterization of a functional Nf-Kappa B site in the human interleukin 1 beta promoter: evidence for a positive autoregulatory loop. *Mol Cell Biol.* (1993) 13:6231–40. doi: 10.1128/mcb.13.10.6231-6240.1993

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

**Publisher’s Note:** All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2022 Kim, Lee, Kim, Park, Han, Kim, Kim and Choi. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.