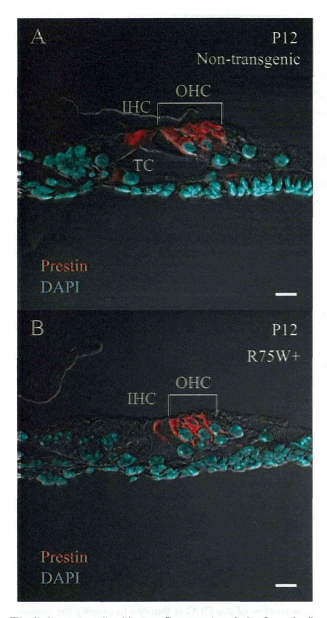
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Hig. 44. A cross-sectional immunofluorescent analysis of prestin distributed in the apical turns of the cochlea of non-transgenic (A) and R75W transgenic mice (B) at P12. Prestin labeling (red) is clearly visible om the whole OHC basolateral wall in both the non-transgenic (A) and R75W transgenic mice (B)) at P12. The extracellular space around the OHC in R75W transgenic mice is narrower than that in non-transgenic mice. On the other hand, the nucleus stained with DAR1 (blue) and the cuticular plate of both mice are devoid of immunostaining. Abbreviations used: OHC, outer hair cell; IHC, inner hair cell. Scale bars are 10 μm (A, B).

arising from the failure of development of the supporting cells can be proposed. First, mature OHCs are supported by underlying Deiter's cells, flanked on the lateral edge by a several rows off Hensen's cells, and anchored by the reticular lamina at their apical surface. The three-dimensional structure off the OHCs enable the longitudinal changes driven by thansmembrane potential changes. In

the transgenic mouse, the OHCs were compressed by the surrounding Deiter's cells, thus restricting motility. Second vibration of the basilar membrane may be related to its thickness, which would contribute to the sensitivity and the production of the otoacoustic emissions (Kossl and Water. 1985) and further to the tonotopic changes of the developing gerbil cochlea (Schweitzer et al., 1996). The thickened basilar membrane observed in the transgenic mice might suppress the DPOAE by reducing the basilar membrane vibration. Structural changes in the basilar membrane may also reduce the sound-induced vibration off the cochlear partition, thus inhibiting deflection of stereocilia on inner hair cells. This could explain why Gib2 R75W/ transgenic mice show remarkable elevation of the auditory brainstem response threshold (Inoshita et al., 2008). Third. morphometric analysis of the organ of Conti suggest possible changes in ionic composition of the contilymph surrounding the basolateral surface of the OHCs (Imposhita et al., 2008). Increased K+ ions in the cortilymph would de-

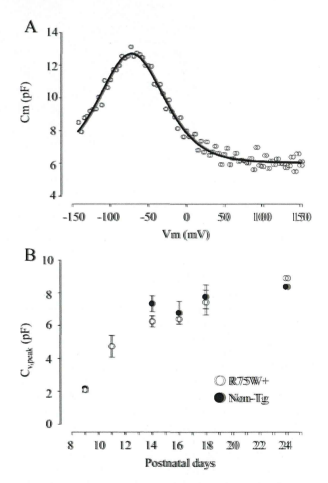


Fig. 5. Electrical responses of isolated OHC. $C_{\rm fm}$ is expressed as a function of $V_{\rm m}$ at P14 in the R75W transgenic mouse (A)). Fitted parameters are $Q_{\rm max}=0.704$ pC, z=0.89. $C_{\rm c}$, peak is expressed as a function of postnatal day (B). The number of cells in non-transgenic (closed circle) and R75W transgenic mice (open circle) was (from P9) to P24) 1–2, 0–3, 2–3, 5–2, 3–3, and 1–1, respectively. Standarderow is plotted. Non-Tg: non-transgenic mice, R75W+: R75W/ transgenic mice.

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polarize the OHCs, and decreased driving force across the mechanosensitive channels could affect OHC electromotility. The progressive degeneration of OHCs observed in the adult R75W transgenic mice (Kudo et al., 2003) may be brought about by disturbed homeostasis of the contilymph.

The secondary hair cell loss in adult R75W transgenic mice (Kudo et al., 2003; Inoshita et al., 2008) implies that the restoration of hearing requires the regeneration of hair cells in addition to introduction of the Gib2 gene. The present study clearly showed both morphological and functional maturation of OHC until late in development, suggesting that a dominant-negative R75W mutation of Gjb2 does not affect the genes that determine or control the differentiation of the OHC. Therefore, gene transfer of Gjb2 into the supporting cells before hair cell degeneration could be used to treat deafness. Transgene expression has been accomplished in the supporting cells of the neonatal mouse cochlea using adeno-associated viral vectors without causing additional damage to the cochlea (lizuka et al., 2008). Therefore, the present study provides a new strategy to restore hearing in Gjb2-based mutation.

CONCLUSION

OHIC from the dominant-negative R75W mutation of *Gjb2* showed normal development and maturation, and isolated OHIC clearly showed voltage-dependent, nonlinear capacitance with characteristic subcellular features. However, the DPOAE, which serves as an index for *in vivo* cochlear amplification, was remarkably suppressed in the mutant mice. This may result from disturbed development of the supporting cells surrounding the OHCs. The present study confirmed that the normal development of the supporting cells is indispensable for the cellular function off the OHIC.

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←前号に続く

3. Cell therapy targeting cochlear fibrocytes

神谷 和作 順天堂大学医学部耳鼻咽喉科学教室

Cell therapy targeting cochlear fibrocytes

Kazusaku Kamiya Juntendo University School of Medicine, Department of Otoralyngology

Recently, a number of clinical studies for cell therapy have been reported and clinically used for several intractable diseases. Inner ear cell therapy for sensorineural hearing loss also has been studied using some laboratory animals, although the successful reports for the hearing recovery were still few.

Cochlear fibrocytes play important roles in normal hearing as well as in several types of sensorineural hearing loss due to inner ear homeostasis disorders. Recently, we developed a novel rat model of acute sensorineural hearing loss due to fibrocyte dysfunction induced by a mitochondrial toxin^{1], 2]}. In this model, we demonstrate active regeneration of the cochlear fibrocytes after severe focal apoptosis without any changes in the organ of Corti. To rescue the residual hearing loss, we transplanted mesenchymal stem cells into the lateral semicircular canal; a number of these stem cells were then detected in the injured area in the lateral wall. Rats with transplanted mesenchymal stem cells in the lateral wall demonstrated a significantly higher hearing recovery ratio than controls. The mesenchymal stem cells in the lateral wall also showed connexin 26 and connexin 30 immunostaining reminiscent of gap junctions between neighboring cells³⁾. These results indicate that reorganization of the cochlear fibrocytes leads to hearing recovery after acute sensorineural hearing loss in this model and suggest that mesenchymal stem cell transplantation into the inner ear may be a promising therapy for patients with sensorineural hearing loss due to degeneration of cochlear fibrocytes.

Key words: cochlear fibrocyte, inner ear cell therapy, mesenchymal stem cell 和文キーワード: 蝸牛線維細胞, 内耳細胞療法, 買業系幹細胞

Mammalian cochlear fibrocytes of the mesenchymal nonsensory regions play important roles in the cochlear physiology of hearing, including the transport of potassium ions to generate an endocochlear potential in the endolymph that is essential for the transduction of sound by hair cells^{4), 5), 6)}. It has been postulated that a potassium recycling pathway toward the stria vascularis via fibrocytes in the cochlear lateral wall is critical for proper hearing, although the exact mechanism has not been definitively proven⁵⁾. One candidate model for this ion transport system consists of an extracellular flow of potassium ions through the scala

tympani and scala vestibuli and a transcellular flow through the organ of Corti, supporting cells, and cells of the lateral wall^{70, 80}. The fibrocytes within the cochlear lateral wall are divided into type I to V based on their structural features, immunostaining patterns, and general location⁸⁰. Type II, type IV, and type V fibrocytes resorb potassium ions from the surrounding perilymph and from outer sulcus cells via the Na, K-ATPase. The potassium ions are then transported to type I fibrocytes, strial basal cells and intermediate cells through gap junctions, and are secreted into the intrastrial space through potassium channels. The

secreted potassium ions are incorporated into marginal cells by the Na, K-ATPase and the Na-K-Cl cotransporter, and are finally secreted into the endolymph through potassium channels.

Degeneration and alteration of the cochlear fibrocytes have been reported to cause hearing loss without any other changes in the cochlea in the Pit-Oct-Unc (POU)-domain transcription factor Brain-4 (Brn-4) deficient mouse⁹⁾ and the otospiralin deficient mouse⁶⁾. Brn-4 is the gene responsible for human DFN3 (Deafness 3), an X chromosome-linked nonsyndromic hearing loss. Mice deficient in Brn-4 exhibit reduced endocochlear potential and hearing loss and show severe ultrastructural alterations, including cellular atrophy and a reduction in the number of mitochondria, exclusively in spiral ligament fibrocytes 9). 10). In the otospiralin deficient mouse, degeneration of type III and IV fibrocytes is the main pathological change and hair cells and the stria vascularis appear normal⁶⁾. Furthermore, in mouse and gerbil models of age-related hearing loss 111), 12), 13), degeneration of the cochlear fibrocytes preceded the degeneration of other types of cells within the cochlea, with notable pathological changes seen especially in type II, IV, and V fibrocytes. In humans, mutations in the connexin 26 (Cx26) and connexin 30 (Cx30) genes, which encode gap junction proteins and are expressed in cochlear fibrocytes and non-sensory epithelial cells, are well known to be responsible for hereditary sensorineural deafness. 140, 150. These instances of deafness related to genetic, structural and functional alterations in the cochlear fibrocytes highlight the functional importance of these fibrocytes in maintaining normal hearing.

Generation of the animal model to study cochlear fibrocyte

To study the role of cochlear fibrocytes in hearing loss and hearing recovery, we developed an animal model of acute sensorineural hearing loss due to acute cochlear energy failure by administering the mitochondrial toxin 3-nitropropionic acid (3NP) into the rat round window niche^{101, 20}. 3NP is an irreversible inhibitor of succinate dehydrogenase, a complex II enzyme of the mitochondrial electron transport chain^{160, 170}. Systemic administration of 3NP has been used to produce selective striatal degeneration in the brain of several mammals^{180, 190}. Our model with 3NP administration into the rat cochlea showed acute sensorineural hearing loss and revealed an initial pathological change in the fibrocytes of the lateral wall and spi-

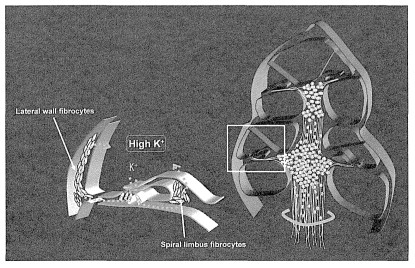


Figure. 1

The localization and the function of cochlear fibrocytes. In mammalian cochlea, ATP-dependent potassium recycling pathways have been well known as the essential mechanism for normal sound input. Cochlear fibrocytes in lateral wall and spiral limbus play a critical role in this potassium recycling system. They transport K+ into the endolymph and keep high K+ concentration mainly by Na+/K+-ATPase and gap innetion.

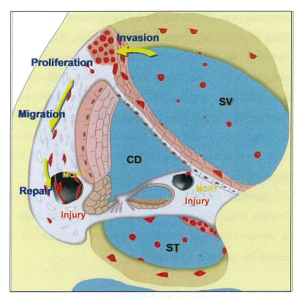


Figure. 2

A summary of the histological observations and our hypothesis for the migration of the transplanted MSCs. Arrows indicate the hypothetical route of MSC migration to the injured area. Some MSCs formed a cell mass around the scala tympanii. A number of MSCs successfully invaded the lateral wall. The invading MSCs migrated and proliferated in the lateral wall. Cell migration may be induced by some chemokines such as MCP1 which was detedted in our DNA microarray analysis. The MSCs which reached the injured area continued to proliferate and repaired the disconnected gap junction network. SW, scala vestibuli; CD, cochlear duct; ST, scala tympanii. The schematic illustration was cited and modified from Am J Pathol, 1711: 214-226, 2007 Kamiya, et al.

ral limbus without any significant damage to the organ of Corti or spiral ganglion. Furthermore, depending on the dose of 3NP used, these hearing loss model rats exhibited either a permanent threshold shift (PTS) or a temporary threshold shift (TTS). In the following study, we used doses of 3NP that induce TTS to explore the mechanism of hearing recovery after injury to the cochlear fibrocytes, and examined a novel therapeutic approach to repair the injured area using mesenchymal stem cell (MSC) transplantation.

Mesenchymal Stem Cell (MSC) Transplantation

MSCs are multipotent cells that can be isolated from adult bone marrow and can be induced to differentiate into a variety of tissues in vitro and in vivo²⁰⁾. Human MSCs transplanted into fetal sheep intraperitoneally undergo site-specific differentiation into chondrocytes, adipocytes, myocytes, cardiomyocytes, bone marrow stromal cells, and thymic stroma²¹⁾. Furthermore,

when MSCs were transplanted into postnatal animals, they could engraft and differentiate into several tissue-specific cell types in response to environmental cues provided by different organs²²⁾. These transplantability features of MSCs suggested the possibility that they could restore hearing loss in 3NP-treated rats to the normal range. Recently, experimental bone marrow transplantation into irradiated mice suggested that a part of spiral ligament which consists of cochlear fibrocytes was derived from bone marrow cells or hematopoietic stem cells²³⁾. This indicates that bone marrow derived stem cells such as MSC may have a capacity to repair the injury of cochlear fibrocytes.

MSC transplantation accelerated hearing recover

The 3NP-treated rats showed complete hearing recovery at low frequencies; however, there remained a residual hearing loss at higher frequencies. Considering that the cochlear fibrocytes that were injured in this model are mesenchymal in origin, we transplanted rat MSCs into the cochlea to attempt to rescue the residual hearing loss. We used MSC which we previously established and demonstrated their potential as MSC, and we further confirmed the surface antigen expression of the cells used for transplantation in flow cytometry which showed similar expression pattern to human and murine MSCs. This suggests that the cells maintained the capacity as rat MSC at the moment of transplantation. Because there is no barrier in the inner ear perilymph between the cochlear and vestibular compartments, cells delivered from the lateral semicircular canal by perilymphatic perfusion are considered to have reached the cochlea. Within the perilymph of the cochlea, these cells presumably spread through the scala vestibuli toward the apical turn of the cochlea, and then, after passing through the helicotrema where the scala vestibuli communicates with the scala tympani, kept moving through the scala tympani toward the basal turn. There is no other way in which MSCs can spread within the cochlear perilymph.

Invasion of MSC to lateral wall tissue

Our study clearly demonstrates that rat MSCs were

successfully transplanted into the inner ear of 3NPtreated rats by perilymphatic perfusion from the lateral semicircular canal. A number of MSCs were detected on the surface of the ampullary crest facing the perilymph and some of them were detected within the tissue of the ampullary crest, indicating that MSCs survived at least for 11 d after the perfusion and had maintained their ability to invade and migrate into the inner ear tissue. In the cochlea, a number of MSCs formed cell masses on the surface of the scala timpanii. where the majority of the surrounding tissue is bone tissue, suggesting that these MSCs did not invade the cochlear tissue. In the scala vestibuli, a small number of MSCs were also found attached to the surface of the bone and the Reissner membrane. However, in the apical part of the lateral wall, a number of MSCs were observed within the tissue, suggesting that MSCs had successfully invaded the lateral wall from the perilymph. This area may be an optimum site for MSC invasion. Furthermore, we performed DNA microarray analysis of the cochlear lateral wall RNAs in 3NPtreated rats and found a significant increase in the expression of the small inducible cytokine A2 gene encoding monocyte chemoattractant protein 1 (MCP1), which has been reported as a chemokine that induces migration of neural stem cells24). This may suggests that the MSC migration to the injured area of the lateral wall in this study may also be induced by chemokines because most MSCs were obsereved in the lateral wall in basal turn which had a prominent damage, but not in the apical turn.

Conclusion

Bone marrow MSCs have greater advantages for clinical use in human subjects than other multipotential stem cells, such as embryonic stem cells, because MSCs can be collected from the patient's own bone marrow for an autologous transplantation with little physical risk, no rejection risk, and few ethical problems. In the present transplantation, many MSCs were confirmed to have invaded the lateral wall and to have contributed to recovery of hearing loss despite transplantation between different rat strains. Therefore, we expect that autologous transplantation

of bone marrow MSCs would be even more effective in treating hearing loss caused by injuries to the cochlear fibrocytes. In addition, significant improvement of hearing by MSC transplantation between different rat strains indicates a possibility of allogenic transplant. Even temporary effects by allogenic transplant may cause difference in the final outcome of hearing recovery by promoting regeneration or viability of host fibrocytes during acute period of injury.

Cell therapy targeting regeneration of the cochlear fibrocytes may therefore be a powerful strategy to cure sensorineural hearing loss that cannot be reversed by current therapies.

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POSTNATAL DEVELOPMENT OF THE ORGAN OF CORTI IN DOMINANT-NEGATIVE GJB2 TRANSGENIC MICE

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Abstract—Hereditary hearing loss is one of the most prevalent inherited human birth defects, affecting one in 2000. A strikingly high proportion (50%) of congenital bilateral nonsyndromic sensorineural deafness cases have been linked to mutations in the GJB2 coding for the connexin26. It has been hypothesized that gap junctions in the cochlea, especially connexin26, provide an intercellular passage by which K+ are transported to maintain high levels of the endocochlear potential essential for sensory hair cell excitation. We previously reported the generation of a mouse model carrying human connexin26 with R75W mutation (R75W+ mice). The present study attempted to evaluate postnatal development of the organ of Corti in the R75W+ mice. R75W+ mice have never shown auditory brainstem response waveforms throughout postnatal development, indicating the disturbance of auditory organ development. Histological observations at postnatal days (P) 5-14 were characterized by i) absence of tunnel of Corti, Nuel's space, or spaces surrounding the outer hair cells, ii) significantly small numbers of microtubules in inner pillar cells, iii) shortening of height of the organ of Corti, and iv) increase of the cross-sectional area of the cells of the organ of Corti. Thus, morphological observations confirmed that a dominant-negative Gjb2 mutation showed incomplete development of the cochlear supporting cells. On the other hand, the development of the sensory hair cells, at least from P5 to P12, was not affected. The present study suggests that Gib2 is indispensable in the postnatal development of the organ of Corti and normal hearing. © 2008 IBRO. Published by Elsevier Ltd. All rights reserved.

Key words: hereditary hearing loss, mouse, organ of Corti, Gjb2.

Hereditary deafness affects about one in 2000 children and mutations in the connexin26 (Cx26) gene (*GJB2*) are the most common genetic cause of congenital bilateral non-syndromic sensorineural hearing loss. It has been

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hypothesized that gap junctions in the cochlea, especially Cx26, provide an intercellular passage by which K+ are transported to maintain high levels of the endocochlear potential (EP), which is essential for sensory hair cell excitation. However, the pathogenesis of deafness remains unresolved because the electrophysiological and histological examination that can be carried out in humans is limited and partly because Gjb2 deficient mice were embryonic lethal (Gabriel et al., 1998). We previously reported om transgenic mice (Tg) carrying human Cx26 with a R7/5W/ mutation that was identified in a deaf family with autosomal dominant negative inheritance. Although the EP nemained within a normal range, the auditory brainstem response (ABR) revealed that the mice at postnatal day 14 (P14) showed severe to profound hearing loss. The tunnel off Conti was not detected and the shapes of outer hair cells (OHCs) were peculiar in the Tg mice at P14. These results suggested that the Gjb2 mutation primarily disturbs homeostasis of cortilymph, an extracellular space surrounding the sensory hair cells, due to impaired potassium ion transport by supporting cells, secondarily resulting in degeneration of the organ of Corti, rather than affecting endolymph homeostasis in mice (Kudo at al., 2003).

Gap junctions are believed to be important for maturation and differentiation of developing tissues (Elias et al., 2007). Developmental expression of Cx26 in the mouse cochlea started in the inner and outer sulcus cells on the 18th day of gestation. At birth, immunolabeling for Cx26 was observed over the supporting cells of the inner hair cells (IHICs) and the mesenchymal components of the stria vascularis (Frenz and Van de Water, 2000). In contrast, Cx26 was not detected in the supporting cells in the organ of Corti before P3. Not until P8 was Cx26 immunoreactivity detected in almost all supporting cells in the organ of Corti ((Zhang et al., 2005)). Thus, evaluation of the postnatal development of Gjb2 Tg mouse cochlea is required to obtain a better and accurate understanding of the molecular mechanism mediated by Gjb2 mutation.

The present study was designed to evaluate the organ of Conti im the R75W+ mice compared with that of non-Tg mice from P5 to P14.

EXPERIMENTAL PROCEDURES

Animals and anesthesia

All mice used throughout this study were obtained from breeding colony, with R75W+ mice (Kudo et al., 2003) and maintained at Institute for Animal Reproduction (Ibaraki, Japan). R75W+ mice were maintained on a mixed C57BL/6 background and intercrossed to generate R75W+ animals. The animals were genotyped using DNA obtained from tail clips and amplified with the

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Abbreviations: ABR, auditory brainstern response; Cx26; connexin26; DC, Deiter's cell; EDTA, ethylenediaminetetraacetic acid; EP; endo-cochlear potential; FGFR3, fibroblast growth factor receptor 3; GA, glutaraldehyde; GJB2, connexin26 gene; HI-E, hematoxylinandieosin; HI-C, inner hair cell; IPC, inner pillar cell; OHC, outer hair cell; OPC, outer pillar cell; P, postnatal day; PB, phosphate buffer; PBS; phosphate-buffered saline; PFA, paraformaldehyde; TIEM, transmission electron microscopy; Tg, transgenic.

Tissue PCR Kit (Sigma, St. Louis, MO, USA). All experiment protocols were approved by the Institutional Animal Care and Use Committee at Juntendo University, and were conducted in accordance with the US National Institutes of Health Guidelines for the Care and Use of Laboratory Animals. We minimized number of animals used and their suffering. Animals were deeply anesthetized with an i.p. injection of ketamine (100 mg/kg) and xylazine (10 mg/kg) in both ABR measurements and histological examinations.

ABR

All electrophysiological examinations were performed within an acoustically and electrically insulated and grounded test room. Mice ranging in age from P10 to P14 were studied. For ABR measurement, stainless-steel needle electrodes were placed at the vertex and ventrolateral to the left and right ears. The ABR was measured using waveform storing and stimulus control of Scope software of Power Laboratory system (model PowerLab4/25, AD Instruments, Castle Hill, Australia), and electroencephalogram recording was made with an extracellular amplifier AC PreAmplifier (model P-55, Astro-Med, West Warwick, RI, USA). Acoustic stimuli were delivered to the mice through a coupler type speaker (model: ES1spc, Bio Research Center, Nagoya, Japan). The threshold was determined for frequencies of 12, 24, 36, and 48 kHz from a set of responses at varying intensities with 5 dB intervals and electrical signals were averaged at 512 repetitions. If the hearing threshold was over 95 dB, it was determined as 100 dB.

Light microscopy

The animals were deeply anesthetized and perfused intracardially with 0.01 M phosphate-buffered saline (PBS; pH 7.2), followed by 4% paraformaldehyde (PFA; pH 7.4) in 0.1 M phosphate buffer (PB; pH 7.4). The mice were decapitated and their cochleae dissected out under a microscope and placed in the same fixative at room temperature for overnight. Cochlear specimens were then placed into 0.12 M EDTA (pH 7.0) in PBS for decalcification for a week, dehydrated and embedded in paraffin. Serial sections (6 μ m) were stained with hematoxylin and eosin (H-E) staining.

Transmission electron microscopy (TEM)

The animals were deeply anesthetized and perfused intracardially with 0.01 M PBS, followed by 4% PFA and 2% glutaraldehyde (GA) in 0.1 M PB. The cochleae were opened and flushed with buffered 4% PFA and 2% GA and fixed for 2 h at room temperature. After washing, the specimens were post-fixed 1.5 h in 2% OsO_4 in 0.1 M PB, then dehydrated through graded ethanols and embedded in Epon. The samples were cut (1 μ m), stained with uranyl acetate and lead citrate, and examined by electron microscopy (H-7100, Hitachi, Tokyo, Japan).

In order to observe microtubules of inner pillar cell (IPC), the mice at the age of P12 were selected. Cochleae were perfused *in situ* with 2.5% GA in 0.1 M PB (pH 7.4) containing 2% tannic acid through the round window, dissected and immersed in the same fixative for 2 h at room temperature. Post-fixation, dehydration and embedding were performed as described above. Ultrathin sections, 60 nm thick, were cut in cross-section.

Immunohistochemistry

The cochleae were removed after cardiac perfusion with 4% PFA, placed in the same fixative at room temperature for an hour, decalcified with 0.12 M EDTA at 4 °C overnight, cryoprotected in 30% sucrose, embedded in OCT, and 10- μ m-thick-sections were collected. Sections were washed in several changes of 0.01 M PBS, blocked with 0.3% Triton X-100 in 0.01 M PBS for 30 min, and then incubated overnight at 4 °C with primary antibody diluted

in 0.01 M PBS+0.3% Triton X-100. The following day, the tissues were rinsed with 0.01 M PBS, incubated for 6 h at 4 °C with a fluorescent-conjugated secondary antibody, rinsed with 0.01 M PBS, and then mounted in Vectashield containing DAPI (Vector Laboratories, Burlingame, CA, USA). The following primary antibodies were used: rabbit polyclonal antibodies to rabbit polyclonal antibodies to fibroblast growth factor receptor 3 (FGFR3) (1:200; Santa Cruz Biotechnology, Santa Cruz, CA, USA), rabbit polyclonal antibodies to p27^{Kip1} (1:200; Lab Vision, Fremont, CA,

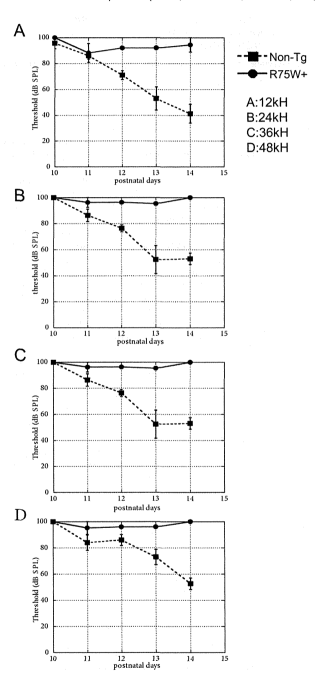


Fig. 1. Developmental change of the threshold levels of ABR of non-Tg and R75W+ mice at 12 kHz (A), 24 kHz (B), 36 kHz (C) and 48 kHz (D). The onset of hearing in non-Tg mice appears at P11, and ABR thresholds achieve adult level (dotted lines in A–D). ABR of R75W+ mice at P11 shows severe to profound deafness at overall sound pressure level (solid lines in A–D).