ILDR1 null mice, a model of human deafness DFNB42, show structural aberrations of tricellular tight junctions and degeneration of auditory hair cells

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In the mammalian inner ear, bicellular and tricellular tight junctions (tTJs) seal the paracellular space between epithelial cells. Tricellulin and immunoglobulin-like (Ig-like) domain containing receptor 1 (ILDR1, also referred to as angulin-2) localize to tTJs of the sensory and non-sensory epithelia in the organ of Corti and vestibular end organs. Recessive mutations of TRIC (DFNB49) encoding tricellulin and ILDR1 (DFNB42) cause human nonsyndromic deafness. However, the pathophysiology of DFNB42 deafness remains unknown. ILDR1 was recently reported to be a lipoprotein receptor mediating the secretion of the fat-stimulated cholecystokinin (CCK) hormone in the small intestine, while ILDR1 in EpH4 mouse mammary epithelial cells in vitro was shown to recruit tricellulin to tTJs. Here we show that two different mouse Ildr1 mutant alleles have early-onset severe deafness associated with a rapid degeneration of cochlear hair cells (HCs) but have a normal endocochlear potential. ILDR1 is not required for recruitment of tricellulin to tTJs in the cochlea in vivo; however, tricellulin becomes mislocalized in the inner ear sensory epithelia of ILDR1 null mice after the first postnatal week. As revealed by freeze-fracture electron microscopy, ILDR1 contributes to the ultrastructure of inner ear tTJs. Taken together, our data provide insight into the pathophysiology of human DFNB42 deafness and demonstrate that ILDR1 is crucial for normal hearing by maintaining the structural and functional integrity of tTJs, which are critical for the survival of auditory neurosensory HCs.

INTRODUCTION

A variety of intercellular structures seal the paracellular space between epithelial cells that outline and separate different cavities and structures in the body. Hemidesmosomes, desmosomes and adherens junctions compose adhesion junctions between neighboring epithelial cells while gap junctions provide an intercellular communication pathway (1). Tight junctions (TJs) between adjacent epithelial cells are ion- and size-selective barriers that regulate the lateral diffusion of solutes and water molecules through the paracellular pathways (2,3). Bicellular tight junctions (bTJs) form a network of parallel, paired strands comprised of integral membrane proteins, membrane-associated and intracellular proteins (3,4). These protein complexes of paired

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TJ strands from two adjacent cells associate laterally with one another to contribute to the semi-permeable seals formed within the paracellular space. On the intracellular side of the plasma membrane, a variety of adaptor proteins are involved in the formation of TJs, such as zona occludens (ZO)-1, ZO-2 and ZO-3, which connect TJs with the actin cytoskeleton, signaling molecules as well as membrane proteins (3,5).

Many integral membrane TJ proteins have been identified, including a family of 24 different claudins (2), junctional adhesion molecules and the TJ-associated MAL and related proteins for vesicle trafficking and membrane link (MARVEL) proteins occludin (6), tricellulin (also known as marvelD2) (7,8) and marvelD3 (3,9). Occludin was the first component of TJs to be identified (6) and has been shown to form heteromeric complexes with tricellulin (10). Claudins play a key role in TJ formation through lateral association and homotypic adhesive interactions between claudins in adjacent cells (3). Claudin-based pores formed between cells in different epithelia show variation in permeability and ionic charge preferences, thus creating a wide range of functionally distinct TJ paracellular barriers throughout the body (2).

In the mammalian cochlea, TJs separate the endolymphatic, perilymphatic and intrastrial fluid spaces, three compartments with distinctly different fluid compositions (11–13). Additionally, TJs play a key role in the generation and maintenance of the endocochlear potential (EP) of 80–120 mV, essential for normal hearing, by sealing epithelial paracellular spaces to separate the potassium-rich endolymph from sodium-rich perilymph (Fig. 1)(14). The bTJs of the reticular lamina have a unique, elaborate structure (11) composed of a fusion of TJs and tight adherens junctions (TAJs) (15). These TAJs are large and, as revealed by freeze-fracture, characterized by strands present along the entire junction depth unlike TJs in most epithelial cells (11).

Mutations of several genes encoding TJ proteins have been found to cause deafness in humans, mice and zebrafish (Table 1). Mutant alleles of human CLDN14, encoding claudin 14, cause nonsyndromic deafness DFNB29 (20). Deaf, claudin-11-deficient mice lack an EP due to impaired TJs between basal cells of the stria vascularis (23,24). In contrast, mutations in Cldn9, encoding claudin 9, and Cldn14 have altered TJs of the reticular lamina but are able to maintain a normal EP (21,22). Hearing loss in both of these mouse models was attributed to leakage of potassium ions through the paracellular pathway of the organ of Corti reticular lamina into the basolateral domain. The increased concentration of potassium in the perilymph surrounding free lateral surfaces of outer hair cells (OHCs) creates a toxic microenvironment, contributing to progressive hair cell (HC) loss in both of these mouse models during the late postnatal period when the EP and hearing are being established in the mouse (21,22).

Tricellular tight junctions (tTJs) serve as sealing elements of the central tube formed when the parallel-paired strands of TJs from three adjacent cells come together and extend basally at the location where three epithelial cells meet (4,26–28). While bTJs are responsible for approximately 99% of the paracellular ion transport, tTJs allow for the passage of various molecules and macromolecules through the tricellular central tube with tricellulin expression regulating the size and selectivity of the pore (26). Tricellulin was the first identified component of tTJs and mutations of TRIC, encoding tricellulin, are associated with human nonsyndromic recessive deafness DFNB49 (7,8).

Immunoglobulin-like domain-containing receptor 1 (ILDR1, also referred to as angulin-2) was recently reported as a lipoprotein receptor mediating the secretion of fat-stimulated cholecystokinin (CCK) hormone in the small intestine (29) and was also identified as a TJ protein (30). Mutations of human ILDR1 cause nonsyndromic deafness DFNB42 (16). In the mouse organ of Corti, ILDR1 localizes to tTJs of both sensory and non-sensory epithelia (30). ILDR1 is a member of the angulin family of TJ-associated proteins that includes angulin-1/lipolysis-stimulated lipoprotein receptor (LSR), angulin-2/ILDR1 and angulin-3/ILDR2 (30,31). Angulin proteins were reported to recruit tricellulin to the tTJs in cultured EpH4 mammary epithelial cells suggesting a dependence of tricellulin on angulin proteins for proper tTJ localization during development (30).

In this study, the effect of loss of ILDR1 (Fig. 2A) on hearing function was characterized using two different functional null mutant alleles of mouse Ildr1:Ildr1GW1(KOMP)Wtsi, a gene trap in intron 2, henceforth abbreviated Ildr1Wtsi (Fig. 2B) and Ildr1m1(kOmp)Wtsi, which deletes exons 3–5 and designated

Figure 1. Tight junction composition and function in the mammalian inner ear. The bTJs and tTJs are composed of integral membrane and membrane-associated proteins that form paired, parallel horizontal and vertical strands with those in adjacent cells (red). In the inner ear, bTJs and TJs help separate K+ rich endolymph from Na+ rich perilymph. Tricellulin (green) and ILDR1 (yellow) localize to tricellular contacts of both sensory and non-sensory epithelia. In the area where three cells meet, tTJ components form a central tube, which is responsible for most macromolecule paracellular flux (orange arrow).
Il2rα−/− (Fig. 2C). As a genomic manipulation can have unexpected off-target effects, the link between a gene and a phenotype is strengthened if the same key features of the phenotype are observed for more than one engineered mutant allele. The hearing phenotype of Il2rα−/− and Il2rκ−/− mice, which are on different genetic backgrounds, was assessed by auditory brainstem response (ABR), distortion product otoacoustic emission (DPOAE) and EP measurements. Furthermore, the inner ear phenotype of Il2rα−/− mice was studied by immunocytochemistry, scanning electron microscopy (SEM), and freeze-fracture electron microscopy.

Whether or not ILDR1 is necessary to recruit tricellulin to tTJs in vivo, specifically in the inner ear, is an unanswered question. The mechanism that targets tricellulin to the tTJs in the cochlea is not well understood and is addressed in our study. Our results indicate that ILDR1 is not necessary for tricellulin recruitment to tTJs in the auditory and vestibular sensory epithelia or in the stria vascularis, but it is essential for retention of tricellulin and the long-term integrity of tTJs. Additionally, we provide insight into potential mechanisms for deafness due to the loss of ILDR1 function.

RESULTS

In wild-type mice, ILDR1 is localized to tTJs of the inner ear

ILDR1 immunoreactivity spans the entire depth of the tTJ as seen in the control Il2rα+/+ organ of Corti labeled with anti-ILDR1 and anti-ZO-1 antibodies (Fig. 3A). TJ-associated protein ZO-1 outlines the OHC cuticular plate in two rings connected by spokes. An upper heart-shape ring defines the apical surface of OHCs, while a lower circular ring corresponds to the shape of the bottom portion of the OHC cuticular plate (Fig. 3A). In wild-type mice, ILDR1 highlights the central tube of tricellular junctions formed by the lateral membranes of the three adjacent cells and is observed along the entire depth of HC tTJs (Fig. 3A). Heterozygous (Il2rα+/−) mice showed staining indistinguishable from Il2rα+/+ mice (data not shown). Thus, in the wild-type mouse inner ear, ILDR1 is present along the entire depth of the organ of Corti tTJs demarcated by two rings of ZO-1 that flank the upper and lower border of the tTJ. ILDR1 was also detected in tTJs of the

Table 1. Hearing loss as an autosomal recessive phenotype associated with mutations of genes encoding TJ proteins

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</tr>
<tr>
<td>Zebrafish</td>
<td>cldnj</td>
<td>Impaired hearing and vestibular function</td>
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*No dominant mutant alleles encoding TJ proteins have been reported to date.

Figure 2. Schematic representation of ILDR1 and the two mutant alleles of mouse Il2r1 used in this study. (A) ILDR1 is a 537 amino acid protein encoded by 8 exons and has an N-terminal signal peptide (red), an Ig-like domain (teal) and a transmembrane domain (orange). To date, three isoforms of ILDR1 have been reported. Exon 6 is alternatively spliced and exon 7 has two donor splice sites (black dotted lines). The epitope for the commercial ILDR1 antibody (Sigma-Aldrich) used in this study, encoded by a portion of exon 7, is shown as green bar above the protein. (B) Gene trap allele of Il2r1Gt(D178D03)Wrst (Il2r1w2/) was produced from insertion of targeting vector (rFlipROSAgeo (Cre)) into intron 2 of Il2r1 (197 bp internal to the exon 2/intron 2 boundary). (C) Il2r1tm1KOMP2Wtsi allele (Il2r1−/−) was generated by targeted replacement of exons 3–5 with a Lac-Z cassette.
vestibular sensory epithelia (Fig. 3B) and in marginal cells of the stria vascularis (Fig. 3C), as well as in other epithelial cells of the inner ear of Ildr1w1 mice (data not shown).

Ildr1w2/2 mice lack ILDR1 at tTJs in the inner ear

The Ildr1w−/− allele was developed previously to investigate the function of ILDR1 in the small intestine (29). Since the inner ear of Ildr1w−/− mice had not previously been examined for ILDR1 localization, we used anti-ILDR1 antibodies to study its localizations in the inner ear sensory epithelium. These mice were also used to examine the specificity of the antiserum against ILDR1. We found that Ildr1w−/− mice have no detectable ILDR1 in tTJs of cochlear and vestibular (saccule, ampules and utricle) sensory epithelia, marginal cells of stria vascularis (Fig. 3D–F) and other epithelial cells of the inner ear (data not shown).

Ildr1w−/− and Ildr1k−/− mice are deaf

To investigate the effect of ILDR1 loss on hearing in Ildr1w−/− mice, we measured ABRs at four frequencies (8, 16, 32 and 40 kHz) (Fig. 4A). Ildr1w−/− mice have severe hearing loss by the second postnatal week, shortly after the onset of hearing in wild-type mice, compared with both Ildr1w+/+ and Ildr1w+/2 littermates. By 8 weeks of age, the hearing loss is profound in homozygous mutant mice (Fig. 4A). In most cases, no measurable ABR responses were detected at the maximum level tested (90 dB peSPL). Ildr1k+/+ and Ildr1k+/2 littermates show elevated thresholds at higher frequencies in an age-dependent manner, which likely is attributed to the background strain (Swiss Webster, Charles River Laboratories International Inc.) of these mice (33). However, thresholds from Ildr1w+/+ and Ildr1w+/− littermates were still markedly lower than that of Ildr1w−/− mice (Supplementary Material, Fig. S1A). These results indicate that hearing loss in Ildr1w−/− mice is an early-onset recessive trait that is rapidly progressive, and severe.

Responses to tone pips (3–42 kHz) and click stimuli were measured in Ildr1k−/− mice of each genotype aged 2, 4 and 8 weeks (Fig. 4D). No ABRs were detected for any of the test stimuli for Ildr1k−/− mice at the ages tested (Fig. 4D). For these mice, threshold was arbitrarily assigned to be 95 dB SPL. At each age, the mean thresholds for Ildr1k+/+ mice were comparable to that of Ildr1k+/− mice across all frequencies.

Figure 3. ILDR1 localizes to tTJs between epithelial cells of the inner ear and is absent in Ildr1w−/− mice. In wild-type mice, (A) 3D reconstruction image from a confocal z-stack shows ILDR1 (green) localization along the entire depth of the tTJ between HCs and supporting cells, connecting the circumferential rings of ZO-1 (red) at the top and bottom of these junctions (a’–a’’). Localization of ILDR1 at tTJs between (B) HCs and supporting cells of vestibular sensory epithelia and (C) between stria vasculars marginal cells of Ildr1w+/+ mice. No ILDR1 was detected in Ildr1w−/− mice (D–F). Asterisks in (A) and (D) indicate magnified HCs. Arrowheads in (B) and (C) point to ILDR1 localization at tricellular contacts. Scale bars: 10 μm.
Ildr1w mice are severely hearing impaired but have a normal EP. (A) Mean ABR thresholds of Ildr1w–/– and Ildr1w+/+ littermates at 2 weeks (n = 8 each), 4 weeks (n = 11 each) and 8 weeks of age (n = 11 each). Ildr1w–/– mice have higher ABR thresholds than those of Ildr1w+/+ littermates starting at 2 weeks, which increased into adulthood. Ildr1w–/– mice have a hearing phenotype indistinguishable from Ildr1k+/- littermates (Supplementary Material, Fig. S1). (B) Mean DPOAE levels for the same Ildr1w–/– and Ildr1w+/+ littermates tested with ABRs. Ildr1w–/– mice have no detectable DPOAE levels at 2, 4 and 8 weeks of age (n = 8, 11, and 11, respectively, per genotype). Therefore, hearing loss in Ildr1w mice by the second postnatal week may be due to variation in the genetic background. (C) EP measurements from adult (P56-85) and P10 Ildr1w–/– and Ildr1w+/+ littermates. P10 Ildr1w–/– mice establish normal EPs under both normoxic and anoxic conditions (n = 6 and 7 for Ildr1w–/– and Ildr1w+/+, respectively). However, the negative EP under anoxic conditions was smaller in adult Ildr1w–/– mice compared with Ildr1w+/+ mice (Student’s t-test, P < 0.05, n = 5, Ildr1w–/– and Ildr1w+/+). (D) Mean ABR thresholds of Ildr1w–/– and Ildr1w+/+ littermates at 2 weeks (n = 7 each), 4 weeks (n = 5 Ildr1w–/–, 6 Ildr1w+/+), and 8 weeks of age (n = 8 Ildr1w–/–, 12 Ildr1w+/+) Ildr1w–/– mice showed no responses at 95 dB SPL, the maximum sound level used. Difference in ABR threshold elevation between Ildr1w–/– and Ildr1w+/+ mice by the second postnatal week may be due to variation in the genetic background. Ildr1w–/– and Ildr1w+/+ are on Swiss Webster and C57BL/6N genetic backgrounds, respectively. Mutant alleles affecting hearing on a C57BL/6 background have a more severe deafness phenotype (32). (E) Positive EP and negative EP measurements made in P14 Ildr1w–/–, Ildr1w+/+ and Ildr1k+/- mice. No significant difference was found for positive EP measurements (ANOVA, P > 0.05). However, the anoxic EP showed a significantly lower magnitude in the Ildr1w–/– compared with the Ildr1w+/+ and Ildr1k+/- controls (ANOVA, P < 0.01). Positive EP measurements made in 10 week old Ildr1w–/–, Ildr1w+/+ and Ildr1k+/- mice showed no significant difference (ANOVA, P > 0.05). Error bars (A–E) are shown as ± SD.

Ildr1w–/– and Ildr1w+/– mice have a normal EP

Alterations in TJ barriers of the inner ear may affect the EP. The EP is the transepithelial voltage across the epithelial barrier in the cochlea. A positive EP with a magnitude of 80–100 mV, variation dependent on strain or species, is required for normal hearing. The positive EP is generated by the stria vascularis and depends on intact epithelial barriers, normoxic metabolism and normal ionic conductances in the stria vascularis (13). Furthermore, under acute anoxic conditions, cochlear epithelia generate a negative EP with a magnitude of 30–40 mV. The negative EP depends on intact epithelial barriers and normal ionic conductances, primarily the apical transduction channel.
and the basolateral potassium channels in OHCs (34). Thus, measurements of the positive and negative EP can reveal a variety of defects in the cochlea. We first evaluated the positive EP under normoxic conditions when an anesthetized animal is maintained under normal tissue oxygen levels by spontaneous breathing and the negative EP under anoxic conditions when tissue oxygen levels were depleted due to cessation of spontaneous breathing (Fig. 4C and E). Evaluations were performed in young (P10) and adult (P56-85) Ilดร1w mice and Ilดร1k littermates (Fig. 4C). To investigate if the loss of ILDR1 affects the development of the EP, we tested Ilดร1w mice around the time the EP is established (P10). At this age, there was no significant difference in either the positive EP between Ilดร1w/+/+ (39 ± 11 mV, n = 6) and Ilดร1w−/− (38 ± 13 mV, n = 7) mice or the negative EP (−26 ± 9 mV, n = 6–20 ± 10 mV, n = 7) for Ilดร1w/+/+ and Ilดร1w−/−, respectively, (Fig. 4C), indicating that at this age the epithelial barriers of the inner ear, including the organ of Corti and stria vascula- ria as well as apical and basolateral conductances of the OHCs, were apparently intact. Furthermore, adult Ilดร1w−/− mice had a positive EP of normal magnitude (100 ± 12 mV, n = 7) compared with Ilدرك+/+ littermates (91 ± 14 mV, n = 7, Fig. 4C), indicating that the barrier function of the cochlear epithelia was not impaired and that stria vascularia was still functional. However, the negative EP in adult Ilدرك−/− mice was significantly smaller (−13 ± 5 mV, n = 4), Student’s t-test, P < 0.05) when compared with Ilدرك+/+ mice (−35 ± 15 mV, n = 5) (Fig. 4C) (35, 36). These data indicate that lack of ILDR1 does not cause a delay in establishing the EP or a compromise of the stria vascularia in maintaining the positive EP. Diminution of the negative EP, however, could be the consequence of degeneration of the organ of Corti that results from the lack of ILDR1.

EP was also recorded in Ilدركk mice at P14 and 10 weeks of age (Fig. 4E). At P14, there were no significant differences (analysis of variance (ANOVA), P > 0.05) in the magnitude of the positive EP between Ilدركk+/+ (100 ± 3 mV, n = 4), Ilدركk+/− (97 ± 4 mV, n = 6) or Ilدركk−/− (102 ± 3 mV, n = 4) mice (Fig. 4E). Similarly, the magnitude of the positive EP in 10 week old mice was not significantly altered (ANOVA, P > 0.05) in Ilدرك+/+ (117 ± 6 mV, n = 4), Ilدرك+/− (114 ± 2 mV, n = 4) or Ilدرك−/− mice (114 ± 8 mV, n = 7). However, the negative EP recorded under anoxic conditions in P14 Ilدرك−/− mice (−5 ± 1 mV, n = 4) was significantly smaller (ANOVA, P < 0.01) than that recorded in Ilدرك+/+ (−35 ± 7 mV, n = 4) and Ilدرك+/− (−26 ± 16 mV, n = 6) mice (Fig. 4E). Taken together, the normal magnitude of the positive EP in Ilدرك−/− mice strengthens the conclusion that ILDR1 is neither essential for establishing nor for maintaining a normal positive EP.

ILDR1-deficient mice exhibit rapid base-to-apex OHC degeneration

To investigate the inner ear morphology of ILDR1 null mice, we immunostained the sensory epithelium using myosin VIIa antiserum as a HC marker (37). In control wild-type mice at 2 months of age, there is no OHC loss in the apical and middle turns of the organ of Corti (Fig. 5A), although some loss of OHCs is detected in the basal turn, which likely reflects the Swiss Webster genetic background (Fig. 5A, right panel). In Ilدرك−/− mice, no loss of OHCs is observed at P11 (Fig. 5B). However, degeneration of OHCs is detected at P12 (data not shown) and becomes prominent at P13-P15 (Fig. 5C and D). OHC demise occurs in a mosaic pattern and progresses rapidly until there is a complete loss of OHCs by 1 month of age (Supplementary Material, Fig. S2A–C). Ilدرك+/+ and Ilدرك−/− mice did not show signs of OHC degeneration at this time (Supplementary Material, Fig. S2A and B).

Compared with OHCs, inner hair cell (IHC) degeneration progressed more slowly with pronounced cell death at 2 months of age (Fig. 5B–E). The early-onset and severe hearing loss in the Ilدرك−/− mouse coincides with the degeneration of OHCs (37). Similarly, by P14, Ilدرك−/− OHCs are degenerating with some OHCs remaining in the apical 20% of the cochlear duct (data not shown). In contrast, Ilدرك−/− vestibular HCs at 2 months of age are intact as revealed by confocal microscopy (Supplementary Material, Fig. S2D and E). Consistent with these observations, we did not notice any obvious vestibular dysfunction in affected DFNB42 individuals (16) and no head tossing or circling in Ilدرك−/− mice, although vestibular evoked potentials (VSEPs) were not tested in these mice (38).

Abnormalities of stereocilia in Ilدرك−/− mice portend OHC loss

Using SEM analysis, we show the normal appearance of apices of IHCs and OHCs of control P15 Ilدرك+/+ mice (Fig. 6A–C). In P13 Ilدرك−/− mouse organ of Corti, IHC stereocilia bundles also do not have any obvious abnormalities, while OHCs show a mosaic pattern of degeneration and loss. We observed more pronounced loss of OHCs at the basal and middle turns of the cochlea of P13 Ilدرك−/− mouse (n = 4) compared with the apical turn (Fig. 6D and F), which is consistent with the immunocytochemistry data (Fig. 5C and D). Moreover, in P13 Ilدرك−/− mice, degenerating OHC stereocilia bundles exhibited partial loss of shorter-row stereocilia and fusion of stereocilia with the apical membrane of HCs (Fig. 6D, E, insets). Several HCs had apical membrane distortions and missing stereocilia bundles (Fig. 6E). Subsequently, areas lacking OHCs in the middle turn of the organ of Corti appeared to be populated only with supporting cells that often have enlarged or misshapen apical surfaces occupying the vacant OHC space thus sealing the reticular lamina (Fig. 6E). These changes to the hair bundles at P13 may be a consequence of prior damage elsewhere in the cell body, with subsequent apoptotic loss of OHCs as the cause of rapid progression to profound hearing loss. Because there is a normal negative EP in P10 Ilدرك−/− mice indicating normal OHC function, we investigated if OHC loss or stereocilia abnormalities could be detected by P11. No obvious stereocilia abnormalities or OHC loss was detected by SEM at P11 in the Ilدرك−/− mouse organ of Corti (Supplementary Material, Fig S3) confirming the myosin VIIa immunostaining results at P11 (Fig. 5B). Our SEM data suggest the importance of ILDR1 for OHC survival in the organ of Corti immediately after the onset of hearing in mouse.

Tricellulin phenotype at tTJs in the absence of ILDR1 in the inner ear of Ilدرك−/− mice

To investigate if the absence of ILDR1 affects recruitment or retention of tricellulin in vivo, we investigated tricellulin localization in the Ilدرك−/− inner ear. Tricellulin has been shown...
to localize to TJs of all the epithelial cells that outline the scala media including the TJs of HCs and supporting cells in the organ of Corti and vestibular system (8). In the organ of Corti, tricellulin spans the entire unique depth of the TJs between HCs and supporting cells (8,19), providing an opportunity to investigate mislocalization of tricellulin by light microscopy. Disruption of tricellulin along the length of TJs in the organ of Corti was evaluated by image analysis in *Ildr1<sup>−/−</sup>* mice aged P2–P14, prior to OHC degeneration observed by SEM and myosin VIIa immunofluorescence staining (Figs 5B and 6D–E and Supplementary Material, Fig. S4). The depth of the tricellulin immunofluorescence signal was measured and compared with the depth of TJs demarcated by ZO-1 immunoreactivity as a ratio of distribution (ZO-1:tricellulin). From P2-5, tricellulin appeared to be normally localized in *Ildr1<sup>−/−</sup>* mice (mean 0.98 ratio of distribution, 4.7% difference) compared with control animals (mean

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**Figure 5.** OHCs of *Ildr1<sup>−/−</sup>* mice rapidly degenerate around hearing onset followed by a slow IHC loss in adult mice. Maximum intensity projections show degeneration of auditory HCs in *Ildr1<sup>−/−</sup>* mice. Hair cell bodies were probed with antibody against myosin VIIa (green) and F-actin was labeled with phalloidin (red). (A) Apical, middle and basal sensory epithelium in adult, 2 month old *Ildr1<sup>+/+</sup>* control mice. (B–E) Apical, middle and basal sensory epithelium from P11 *Ildr1<sup>−/−</sup>*, P13 *Ildr1<sup>−/−</sup>*, P15 *Ildr1<sup>−/−</sup>*, and adult *Ildr1<sup>−/−</sup>* mice indicate that HCs develop normally but OHCs begin to degenerate around hearing onset. (E) In adult, 2 month old *Ildr1<sup>−/−</sup>* mice, gradual loss of IHCs is observed. Scale bar: 10 μm.
1.03 ratio of distribution, Student’s t-test, $P > 0.05$, Fig. 7A and E). However, at $\approx P6–P10$, tricellulin mislocalization was observed in Ildr1w+/2 mice (mean 0.94 versus mean 1.07 ratio of distribution, 13.3% difference, Fig. 7B and F, Student’s t-test, $P < 0.005$). By P11, tricellulin was noticeably absent from the top portion (relative to the apical surface of the HC) of tTJs in Ildr1w+/2 mice (mean 0.84 versus mean 1.06 ratio of distribution, 22.7% difference) and was predominantly localized at a lower point near the base of the tTJs (Student’s t-test, $P < 0.0001$, Fig. 7C and G). In control P14 Ildr1w+/+ mice, localization of tricellulin (Fig. 7D) is indistinguishable from its localization in Ildr1w+/2 heterozygotes (Supplementary Material, Fig. S4A and D). In 1 month old Ildr1w+/2 mice that lack OHCs, tricellulin is present at the newly formed tTJs between three supporting cells (Supplementary Material, Figs S4B, C, E, F, H and I, and S5).

Ultrastructure of tTJs is abnormal in Ildr1w−/− mouse inner ear

The architecture of the outer leaflet of the plasma membrane can be examined by freeze-fracture electron microscopy (Fig. 8). In the wild-type mouse, freeze-fracture analyses of inner ear tTJs show strands running perpendicular to the OHC apical surface at the point where a HC and two supporting cells meet and where presumably tricellulin normally localizes. These strands appear as closely adjacent particles in a line. In the Ildr1-deficient mouse, the point of juncture between three cells is less defined and particles are not closely adjacent (Fig. 8B, arrows). Moreover, in the wild-type, some of strands that form the TAJ around the neck of the cell, run into and appear to join to the perpendicular strand at almost right angles. But in the mutant, there are almost no strands of the TJs that converge with a perpendicular strand. Rather, many of the strands defining the TAJ run parallel to the perpendicular line at the point of juncture of the three cells. These data indicate that Ildr1 may contribute to the ultrastructure of inner ear tTJs.

DISCUSSION

Ildr1w−/− and Ildr1k−/− mice recapitulate human deafness DFNB42

ILDR1 null mice (Ildr1w−/−) were generated to study the function of ILDR1 as a receptor in fat-stimulated CCK secretion due to its similarity with family member angulin-1/LSR, which mediates LDL uptake (29). We found that Ildr1w−/− and Ildr1k−/−...
young mice have profound hearing loss that recapitulates human deafness DFNB42 phenotype due to recessive mutations of ILDR1 (16). In Ildr1\(^{\text{tm}1-2}\) mice, the absence of ILDR1 in cochlear tTJs results in a rapid degeneration of OHCs as early as P12, around the onset of hearing in mice (Fig. 5) (39,40), which coincides with the establishment of the mature EP in the basal turn of the cochlea. The presence of both normal appearing OHCs and IHCs in the organ of Corti and the generation of a normal negative EP in Ildr1 null mice prior to the onset of hearing indicate that the lack of functional ILDR1 in these mice does not affect development of the organ of Corti. Moreover, absence of ILDR1 does not alter recruitment of tricellulin to tTJs or localization of ZO-1 to bTJs of the postnatal inner ear during development before the onset of hearing. Pathological

Figure 7. Tricellulin localizes to tTJs in the absence of ILDR1, prior to degeneration of auditory HCs. (A–D) 3D reconstructions of z-stack confocal images from whole-mount organ of Corti samples, probed with antibodies directed against tricellulin (green) and ZO-1 (red). (A) In P4 Ildr1\(^{\text{tm}1-2}\) mice, tricellulin is visualized along the entire tTJ depth. (E) Cartoon schematic of tricellulin localization. Velocity measurement parameters and graph show that tricellulin is localized normally in P2–P5 Ildr1\(^{\text{tm}1-2}\) mice (mean age = 3.81 ± 1.19) compared with control animals, \(P > 0.05, n = 4\) control and 5 experimental animals. (B) By P7 (mean age of P6–P10 group = 7.54 ± 0.72 SD), some mislocalization begins to be observed in Ildr1\(^{\text{tm}1-2}\) mice. (F) Arrowhead in cartoon depicts mislocalization at P7, \(P < 0.005, n = 5\) animals per group. (C) In P11 Ildr1\(^{\text{tm}1-2}\) mice, mislocalization is apparent due to lack of tricellulin signal toward the top of the tTJ and its concentration near the basal end of the tTJ. (G) Arrowhead in cartoon shows interpreted mislocalization at P11, and velocity measurements and graph for the P11–P14 control group show a clear statistical significant mislocalization of tricellulin, \(P < 0.000001, n = 6\) animals per group. 3D reconstruction of (D–H) Ildr1\(^{\text{tm}1-2}\) control mice show tricellulin localization as outlined in the cartoon. Drawings and arrowheads show interpretation of tricellulin mislocalization in Ildr1\(^{\text{tm}1-2}\) mice as the cuticular plate thickens from P4 to P11 and visual description of Velocity measurements. Insets show high magnification of OHCs from the corresponding age and asterisks indicate HCs depicted in magnified panels. Error bars (E–H) are shown as ± SD. Scale bars 10 μm.
changes do begin to occur after P10, and there is an elevated ABR thresholds in \textit{Ildr1}^{m/m} mice detected by the second postnatal week. The degeneration of OHCs and IHCs, and the eventual mislocalization of tricellulin, together with freeze-fracture analysis, demonstrate that ILDR1 is essential for the structural integrity and functionality of inner ear tTJs as well as for the long-term viability of HCs after the onset of hearing.

Is human deafness due to mutations of \textit{ILDR1} nonsyndromic or syndromic?

ILDR1 was shown to be a receptor that mediates the secretion of CCK (satiety hormone) in the presence of fatty acids or lipoproteins (29). This finding raises the possibility that ILDR1 could also have a receptor function in the inner ear for a presently unknown ligand in addition to serving as a TJ protein. Further experiments are needed to investigate this possibility. Although ILDR1 has a function in the small intestine, ILDR1 null mice are not outwardly or physiologically different from wild type. Neither \textit{Ildr1}^{w/w} nor \textit{Ildr1}^{k/k} mice appear to have a weight-gain phenotype associated with the loss of ILDR1 (Sanger institute mouse portal, www.sanger.ac.uk/mouseportal/) (29). To date there are no other discernable phenotypic abnormalities in these mice other than an auditory dysfunction described herein for both mutant alleles (\textit{Ildr1}^{w/w} and \textit{Ildr1}^{k/k}). However, original investigations of deaf DFNB42 individuals might have overlooked clinically relevant subtle abnormalities because the primary focus during ascertainment was limited to deafness (41). Our initial conclusion that mutations of human \textit{ILDR1} cause exclusively nonsyndromic deafness DFNB42 may have to be revised as we learn more about the physiological consequence of a loss of ILDR1 in mouse (16).

\textbf{ILDR1 is not necessary for generation and establishment of an EP}

Higashi and coauthors demonstrated that the angulin family members are important for maintenance of TJ barrier function in angulin-1/LSR knock-down EpH4 cells as detected by a significantly lower transepithelial electrical resistance (30,31). While these results indicated that a barrier function for ILDR1 in the inner ear was highly likely, the presence of a normal, positive EP provides evidence that loss of ILDR1 is not compromising the epithelial barrier in the cochlea to an extent that would diminish the EP. The generation of the EP depends on a low K\textsuperscript{+} concentration in the intrastrial fluid space, which is maintained by marginal cells of stria vascularis (13). The tTJs between marginal cells express both tricellulin and ILDR1 (8,30). The presence of a normal positive EP provides evidence that a paracellular leakage, if present, does not exceed the capacity of marginal cells to maintain the required low K\textsuperscript{+} concentration in the intrastrial space. For at least 2 months of age, the functional activity of the strial marginal cells, as well as the barrier function of the stria vascularis, remains intact during EP establishment in ILDR1-deficient mice. We also note that there is no obvious mislocalization of tricellulin in the marginal cell tTJs of the stria vascularis of \textit{Ildr1}^{m/m} mice. However, we cannot exclude the possibility of partial tricellulin mislocalization from tTJs of marginal cells. It is also possible that in the stria vascularis, another angulin family member, such as LSR...
(previously reported to be expressed in the stria vascularis), ILDR2 (30,42), or an unidentified tTJ protein may fortify the marginal cell barrier and compensate for the loss of ILDR1 from these TJs.

**ILDR1 is necessary for retention but not recruitment of tricellulin at tTJs in the organ of Corti**

Our data indicate that ILDR1 is not necessary for the recruitment of tricellulin to tTJs during development of the inner ear, unlike in angulin-1/LSR knock-down EpH4 cells (30). However, ILDR1 appears to be essential for maintaining the structure and integrity of the tTJs in the inner ear. Higashi and coauthors provided evidence for the recruitment of tricellulin by both FLAG-tagged ILDR1 and ILDR2 in vitro. Similarly, LSR-FLAG had previously been shown to recruit tricellulin using the same EpH4/LSR knock-down cells, and LSR is localized at tTJs of non-sensory epithelia in the inner ear (30,31). Notably, ectopic expression of some substitution mutations of ILDR1 disrupted ILDR1-driven tricellulin recruitment, indicating that there is likely an interaction between the two proteins. In this regard, our data indicates that ILDR1 is necessary for the retention of tricellulin at tTJs in the organ of Corti, but not for its recruitment. Mislocalization of tricellulin in Ildr1<sup>w<sup>−/−</sup></sup> mice is detected prior to hair bundle disorganization observed at P12. It is possible that ILDR1 interacts with tricellulin to assist in maintaining the structure of the tTJs, but is not crucial for its initial recruitment to tTJs in the inner ear as shown for cells in vitro (30). However, we cannot exclude the possibility that ILDR1 may still be necessary for tricellulin recruitment in the other organs and tissues (30).

Another protein, which is absent in EpH4 cells, may be responsible for the proper localization of tricellulin in the inner ear of Ildr1<sup>−/−</sup> mice. Generally, members of the angulin family do not have overlapping expression patterns and ILDR1 is the main family member expressed in the inner ear (30). Expression data from massively parallel signature sequencing libraries (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GPL3835) of inner ear cDNAs indicate angulin-3/ILDR2 mRNA was not detected in the organ of Corti and vestibular sensory epithelia, but it is present in stria vascularis (42). Additionally, localization of angulin-1/LSR is different from that of ILDR1 in the inner ear (30). Functional compensation for the loss of ILDR1 in Ildr1<sup>−/−</sup> mice does not occur since Ildr1 homozygous mutant mice exhibit profound deafness suggesting a unique function for ILDR1 at tTJs. In a recent report (19), localization of ILDR1 was shown to be unaffected in tricellulin-deficient mutant mice, indicating that tricellulin does not recruit ILDR1 to the tTJs of the inner ear, similar to results obtained in vitro (30).

**Possible mechanisms of DFN42 deafness and HC degeneration in Ildr1<sup>−/−</sup> mice**

The selective movement of ions, such as K<sup>+</sup>, across the reticular lamina of the cochlea, is related to the establishment of the negative EP (43–45). A lack of ILDR1, and subsequent dysfunction or degeneration of OHCs in Ildr1<sup>−/−</sup> mice, presumably disrupts the movement of ions across the endolymph–perilymph barrier. OHC loss has been shown to impair the establishment of an anoxic EP (35). Therefore, OHC degeneration in ILDR1-deficient mice may account for the smaller amplitude of anoxic EP in these mice. However, loss of OHCs alone in ILDR1-deficient mice cannot explain the observed increase in ABR thresholds. OHC degeneration would lead to a loss of the cochlear amplifier and hence only a 40–60 dB increase in threshold (46). It seems likely that, due to loss of ILDR1, there is an additional dysfunction elsewhere in the auditory system.

Hair cell degeneration in Ildr1 homozygous mutant mice may be due to malfunction of intracellular signaling, exposure of HCs to a toxic extracellular microenvironment, or both. The tricellulin knock-in mouse model of human DFN49 deafness also showed an OHC degenerative phenotype in the presence of normal EP, similar to Ildr1<sup>−/−</sup> mice (19). When mutant tricellulin mice were mated with POU3F4-deficient mice that do not establish an EP and have a reduced concentration of K<sup>+</sup> as well as other ions in the endolymph (47), HCs in double mutants deficient both for tricellulin and POU3F4 remained intact. This indicated that HC degeneration in these mice may be due to extracellular factors, such as a toxic microenvironment on the lateral walls of HCs due to the altered permeability of tricellular barriers between hair and supporting cells in the absence of tricellulin. Future experiments will determine if HC degeneration in Ildr1<sup>−/−</sup> and Ildr1<sup>k<sup>−/−</sup></sup> mice is a cell non-autonomous effect associated with tTJ breakdown.

The observations in ILDR1 null mice that OHC were affected earlier than IHC and that OHC in the basal and middle turns were more affected than OHC in the apical turn is consistent with a common pattern of cochlear damage that has been found in association with aging, noise trauma and drug-induced ototoxicity (48–50). Underlying reasons for this pattern may include qualitative differences between IHC and OHC. The lateral plasma membrane of an IHC is closely opposed to supporting cells while OHCs are surrounded by perilymph allowing for prestin-driven contraction and elongation of cell bodies in response to auditory stimuli. We hypothesize that these rapid cyclical contractions of OHC bodies, when tTJs lack ILDR1 and are compromised, permit influx of ions and molecules from the potassium-rich endolymph into perilymph creating a toxic microenvironment that trigger irreversible damage to OHCs. When the tTJ barrier is compromised, increased K<sup>+</sup> exposure to the basolateral surfaces of OHCs in the organ of Corti may lead to prolonged depolarization of OHCs due to inhibition of K<sup>+</sup> efflux from HCs during repolarization, which induces HC death (51). Alternatively, loss of tricellulin from the most apical region of the tTJ in Ildr1<sup>−/−</sup> mice begins approximately at the time the Ca<sup>2+</sup> gradient between endolymph and perilymph is being established, while the formation of the Na<sup>+</sup> and K<sup>+</sup> gradients are nearly complete (52,53). An early sign of degeneration at P6-P10, evidenced by tricellulin mislocalization in tTJ, may result in a leaky paracellular pathway with detrimental consequences for the HCs.

The tTJs are responsible for the majority of molecular and macromolecular flux within the paracellular space (26). While it is possible that alterations of the tTJ sealing elements may increase ion paracellular flux, there may also be an increased permeability to ATP across the scala media. Exposure to high levels of sound has been shown to increase ATP concentration in the endolymph (54). Increased ATP concentration and/or impaired molecular flux through the paracellular pathway may
invoke a downstream mechanism leading to HC death. The ability of the boundaries of the scala media of Ildr1<sup>w<sup>−/−</sup></sup> and Ildr1<sup>+/−</sup> mice to maintain a positive EP suggests that the ionic paracellular permeability is largely functionally intact, and the selective increase of flux of other molecules from endolymph such as, for example, ATP could be a more likely explanation for OHC degeneration.

Vestibular HCs in Ildr1<sup>w<sup>−/−</sup></sup> mice do not degenerate. The vestibular end organs do not possess the same electrochemical gradient that is found in the auditory periphery and the basolateral surfaces of vestibular HCs are not as exposed to the surrounding liquid as compared with OHCs. We note that ultrastructural analyses of freeze-fracture replicas from vestibular epithelium of Ildr1<sup>+/−</sup> mice show structural abnormalities similar to those observed in tricellulin-deficient mice. These data suggest that the alterations of a barrier function of the inner ear tTJs in Ildr1<sup>w<sup>−/−</sup></sup> mice and the mechanism of OHC death may be similar to those observed in tricellulin-deficient mice.

The results of our study consolidate the importance of ILDR1 as a TJ protein necessary for HC survival, and for the integrity of the auditory system. Our data indicate that two TJ proteins, tricellulin and ILDR1, are recruited to tTJs independently of one another. However, once in place, these two proteins may interact to maintain the structural integrity and function of tTJs, which are critical for normal hearing function in human and mouse.

MATERIALS AND METHODS

Mutant alleles of mouse Ildr1

All animals in this study were handled according to either the NIH Animal Care and Use Committee guidelines or the UK Home Office regulations and the UK Animals (Scientific Procedures) Act of 1986 (ASPA) under a UK Home Office license, and the study was approved by the King’s College London Ethical Review Committee. Mice were culled using methods approved under these guidelines to minimize any possibility of suffering.

Two different mutant alleles of mouse Ildr1 were used in this study. Ildr1<sup>w<sup>−/−</sup></sup> mice, abbreviated here as Ildr1<sup>w<sup>−/−</sup></sup> knock-out mice, were generated on a Swiss Webster background in Rodger Liddle’s laboratory at the Duke Neurotransgenics facility using gene-trapped ES cell clone D178D03 TBV-2 (29). The targeting vector rFlipROSA<sub>βgeo</sub> (Cre) was integrated in intron 2 of Ildr1 (197 bp internal to the exon 2/intron 2 boundary). Ildr1 failed to amplify in a reverse transcription polymerase chain reaction (RT-PCR) assay using RNA obtained from homozygous mutant mice. The second mutant allele of Ildr1 used in this study, Ildr1<sup>+/−</sup> (KOMP)Wtsi<sub>2</sub>, designated here as Ildr1<sup>+/−</sup>, was generated and maintained on a C56BL/6N background as part of The Wellcome Trust Sanger Institute Mouse Genetics Projects (project ID: 31366) (55,56). The targeting vector was inserted such that exons 3–5 were deleted, supposedly inactivating the gene and inserting a LacZ cassette containing a strong splice acceptor. A high-throughput phenotyping screen identified a hearing deficit in the homozygous mutants at 14 weeks of age (56), but did not detect any other phenotypic deficits from the wide range of assays carried out (www.sanger.ac.uk/mouseportal). A breeding colony was established using heterozygote crosses to generate the necessary genotypes for experimental work.

Genotyping

The Ildr1<sup>w</sup> mice were genotyped as previously described (29), with some modifications. Primers were combined together in a multiplex PCR consisting of one forward primer: Ildr1 forward (5′CTGTCCCTTGTAGTCAGCATC) located in exon 2, and two reverse primers: Ildr1 reverse (5′GCTGACTTGGATTGTCACAT) located in intron 2, and lacZ reverse (5′CAAGGCCGATTAAGTGGGTAAGC) located inside the β-Geo cassette (Fig. 2B). Together, the two Ildr1 primers yield a PCR 464 bp product using genomic DNA from both wild-type and heterozygote mice as a template. The Ildr1<sup>+</sup> forward and lacZ reverse primers amplify a 1215 bp product in all Ildr1<sup>w<sup>−/−</sup></sup> and Ildr1<sup>+/−</sup> mice. Together, heterozygote genomic DNA samples yield both PCR products in the multiplex PCR while wild-type and homozygous mutant mice only result in their respective PCR products (Supplementary Material, Fig. S6A). Ildr1<sup>+</sup> mice continued to be bred using heterozygote crosses on their Swiss Webster background with no further back-crossing.

To genotype the Ildr1<sup>+</sup> allele, genomic DNA was extracted and used as the template for short-range PCR using the forward primer (5′TAATGCGAGCGGCTTGGGA), and the reverse primer (5′GCGAAGTCCCTCTGTATTG) for the wild-type allele to yield a 233 bp PCR product. The forward primer for the mutant allele was the same as that for the wild type, and the reverse primer was (5′TCTGGGATACGGTATGCCC) to yield a 159 bp PCR product. The presence or absence of the cassette (Fig. 2C) was confirmed by testing the neomycin resistance sequence using the forward primer (5′CAAGATGGATTGCAGCGATTCTC) and the reverse primer (5′GACGAGATCTTGCGTTCGCCC) to produce a 600–700 bp PCR product (Supplementary Material, Fig. S6B).

ABR and DPOAE measurements

Auditory testing of Ildr1<sup>w<sup>−/−</sup></sup> and Ildr1<sup>+</sup> mice and their respective littermate controls was completed in different laboratories. Ildr1<sup>w<sup>−/−</sup></sup>, Ildr1<sup>+/−</sup> and Ildr1<sup>+/+</sup> littermates were evaluated at the Mouse Auditory Testing Core Facility, NIDCD. Mice were analyzed and tested in a serial, non-terminal fashion at three ages: 2, 4 and 8 weeks of age. Mice were anesthetized via an intraperitoneal (IP) injection of a solution of Ketamine (56 mg/kg) and Dexamethazone (Pfizer) (0.375 mg/kg). Pre-weaned mice received the required dosage diluted 1:1 in a warm saline solution to have a reasonable volume to inject. All mice were tested inside a sound-proof booth (Acoustic Systems) while resting on a heating pad connected to a temperature controller and rectal probe to maintain body temperature near 37°C.

Both DPOAEs and ABRs were measured in the right ear only using Tucker-Davis Technologies (TDT; Alachua, FL, USA) hardware (RZ6 Multi I/O processor, MF-1 speakers) and software (BioSigRz, v. 5.1). DPOAEs were recorded first using two TDT MF-1 speakers and an ER-10B+ microphone (Etymotic, Elk Grove Village, IL) coupled to the mouse’s ear using a modified pipette tip (a 10 μl pipette tip was trimmed such that length = 10 mm and the tip opening = 2 mm).
DPOAE (2f1 − f2) levels were obtained in response to two primary tones at f1 = 65 dB SPL and f2 = 55 dB SPL with f2 varied between 4–44.8 kHz (5 pts/octave) and f2/f1 = 1.25. The mean noise floor was calculated from three points sampled above and three points sampled below the 2f1 − f2 frequency. Output data presented in dB V were converted to dB SPL offline based on the ER-10B+ microphone’s calibration voltage (dB SPL = 20*log10(V/0.02)+93.9).

ABR thresholds were determined by presenting Blackman-gated tone-burst stimuli (3 ms duration, 29.9/s, alternating polarity) at four frequencies (8, 16, 32, and 40 kHz). Subdermal needle electrodes were placed at the vertex and beneath each pinna, with the non-test (left) ear serving as the ground. Stimuli were delivered via a TDT MF-1 speaker in a closed-field configuration (PVC tubing provided with the speaker terminating in a modified pipette tip of the same dimensions as the one used with the DPOAE microphone). Responses were amplified (20 ×), filtered (0.3–3 kHz) and digitized (25 kHz). Thresholds were determined through visual inspection of stacked waveforms (average of 512–1024 artifact-free responses per waveform).

The stimulus initially was presented at 80 dB peSPL and then decreased in 10 dB steps until the waveform disappeared. The stimulus was then increased and decreased in 5 dB steps until the lowest stimulus level that produced a repeatable waveform (ABR threshold) was determined. At least two waveforms (1024 responses each) were obtained for stimulus levels at and below the lowest stimulus level that produced a repeatable waveform (ABR threshold) was determined. When no repeatable waves were visible at the highest stimulus level tested (90 dB peSPL), the threshold was arbitrarily determined through visual inspection of stacked waveforms by identifying the lowest stimulus level with a recognizable waveform.

EP measurements

EP measurements for Ifdrt7+/− mice and wild-type littermates were conducted in the Section of Molecular Biology and Genetics, NIDCD. Mice were anesthetized with tribromoethanol (Sigma-Aldrich) at a dose of 0.35 mg/g body weight per mouse. EP measurements were made using glass microelectrodes in adult (n = 7 normoxic, n = 5 Ifdrt7+/+, 4 Ifdrt7+/−/− and P10 mice (n = 6 Ifdrt7+/+, n = 7 Ifdrt7+/−/−). Glass microelectrodes, consisted of tips of glass pipettes and silver/chloride electrodes, were prepared as described (58) using a P-97 Flaming/Brown micropipette puller (Sutter Instrument, Novato, CA, USA). Procedures were developed by modifying previously described protocols (53, 59, 60). Measurements were made in the basal turn of the cochlea by a round-window approach through the basilar membrane of the first turn. Anoxia was induced by intramuscular injection of succinylcholine chloride (0.1 μg/g body weight per mouse) after establishment of deep anesthesia followed by additional injection of tribromoethanol (n = 5, 4 for adult mice, n = 6, 7 for P10 mice). Data were recorded digitally (Digidata 1440A and Axoscope 10; Molecular Devices, Sunnyvale, CA, USA) and analyzed using Clampfit 10 (Molecular Devices).

EP measurements in Ifdrt7−/− mice were done at the Wolfson Centre for Age-Related Diseases, King’s College London. Mice were anesthetized with an IP injection of urethane (0.1 ml/10 g of a 20% solution). Negative, anoxia EP, in addition to the normal positive EP, was measured in mice aged 2 weeks (P14, n = 4 Ifdrt7−/−, n = 6 Ifdrt7+/−/− and n = 4 Ifdrt7+/+). Positive, normoxic EP was measured in mice aged 10 weeks (P67–P73, n = 7 Ifdrt7−/−/−, n = 4 Ifdrt7+/−/− and n = 4 Ifdrt7+/+). A tracheal cannula was inserted into the left bulla to reveal the wall of the cochlea and a small hole was made over the basal turn of the cochlea. The tip of a 150 mM KCl-filled glass microelectrode was then inserted into the scala media. The EP of Ifdrt7−/+−, Ifdrt7+/−/− and Ifdrt7+/+ mice was measured as a stable positive potential, referenced to an Ag–AgCl pellet placed under the dorsal skin of the neck (34). Once the stable resting EP had been recorded, the electrode was left in situ and the mouse injected with an overdose of urethane. Following onset of anoxia, the potential fell rapidly and the maximally negative (anoxia) EP was measured (34).

Immunocytochemistry and tricellulin mislocalization analyses

To evaluate morphological changes in the organ of Corti of Ifdrt7−/− mice, after measurement of EP at P14, inner ears from homozygous and control littermates were fixed with 4% paraformaldehyde (PFA) for 2 h at room temperature (RT). After rinsing with phosphate buffered saline (PBS), the inner ears were decalcified with 10% EDTA for 4 h at room temperature (RT). After rinsing with phosphate buffered saline (PBS), the inner ears were decalcified with 10% EDTA for 4 h at RT and then dissected to expose the organ of Corti. In order to visualize the stereocilia bundles, the samples were stained with Rhodamine phalloidin (Life Technologies) diluted in 0.05% Triton-X/PBS for 40 min and after rinsing with PBS mounted in ProLong Gold antifade media (Life Technologies). Specimens were imaged using a Zeiss LSM 710 confocal microscope. Each organ of Corti was examined at 10% intervals along the length of the cochlear duct from the base.

To visualize possible HC degeneration and other morphological alterations in the organ of Corti of Ifdrt7−/−/− mice, cochleae were extracted and fixed overnight at 4°C in 4% PFA diluted in 1× PBS at various ages. Organ of Corti and vestibular sensory epithelia were then microdissected in 1× PBS. The tectorial membrane was removed to expose the sensory
The inner ears of P8 Ildr1w mice with 2.5% glutaraldehyde in 0.1M cacodylate buffer with the apex of cochlear capsule. Cochleae were gently perfused cleaned, membranes removed and a small hole was made in the round and oval windows at the base of the cochlea were dissected out in Leibowitz media (L-15, Invitrogen). The removed from animals after decapitation, and the cochleae were fixed in 2.5% glutaraldehyde in 0.1M cacodylate buffer (EMS) supplemented with 2 mM CaCl2 for 2 h at RT and dehydrated in graded series of ethanol (EtOH). Specimens were then transferred from 100% EtOH to 100% acetone, placed into metal mesh baskets (Ted Pella Inc., Redding, CA, USA), critical point dried from liquid CO2 (CPD030 Critical point Dryer, BAL-TEC AG, Florida, USA), and finally sputter-coated by 4–5 nm thick platinum using turbo-pumped sputter coater Q150T (Quorum Technologies, UK). Samples were mounted on aluminum studs (Electron Microscopy Sciences, Hatfield, PA, USA) and imaged using a field emission scanning electron microscope (S-4800, Hitachi, Japan).

Freeze-fracture analysis of Ildr1w mice

The inner ears of P8 Ildr1w/+ and Ildr1w−/− littermates were removed from animals after decapitation, and the cochleae were dissected out in Leibowitz media (L-15, Invitrogen). The round and oval windows at the base of the cochlea were cleaned, membranes removed and a small hole was made in the apex of cochlear capsule. Cochleae were gently perfused with 2.5% glutaraldehyde in 0.1 M cacodylate buffer with 3 mM CaCl2 through these openings, followed by immersion in the same fixative overnight at 4°C. The sensory epithelia of the inner ear were microdissected and incubated in 25% glycerol for at least 45 min prior to mounting on support planchettes using a yeast in glycerol paste as adhesive, and freezing in a rapidly stirred mixture of propane/isopentane (4:1) chilled in liquid nitrogen. Freeze-fracture replicas were obtained by routine procedures in a Balzers BAF400D apparatus (61). Replicas were viewed and imaged on a JEOL 1200EXII microscope. Digital images are adjusted for optimal contrast and brightness using Photoshop GS5 (Adobe) and are presented in reverse contrast.

Statistics

Statistical analysis of EP data for Ildr1w−/− mice was performed using a two-tailed analysis Student’s t-test of independent variables and graphed using Excel. For each data set on tricellulin mislocalization, the D’Agostino and Pearson omnibus normality test in GraphPad Prism 5 software (La Jolla, CA) was performed to determine Gaussian distribution. Statistical differences between data were then determined by a two-tailed distribution Student’s t-test of independent variables. EP analysis for Ildr1w mice was performed by ANOVA. A resulting P value of <0.05 was considered to be significant for all statistical tests represented by asterisks in graphs. All graphically represented data are shown as an average ± SD.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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