**Tmprss3 loss of function impairs cochlear inner hair cell Kcnma1 channel membrane expression**

Laurence Molina\(^1,\)\(^\dagger\), Lydie Fasquelle\(^2,3,\)\(^\ddagger\), Régis Nouvian\(^2,3\), Nicolas Salvetat\(^1\), Hamish S. Scott\(^4,5\), Michel Guipponi\(^6\), Franck Molina\(^1\), Jean-Luc Puel\(^2,3\) and Benjamin Delprat\(^2,3,\)\(*\)

\(^1\)SysDiag UMR 3145 CNRS/Bio-Rad, Cap Delta/Parc Euromédecine, 1682 rue de la Valsière, CS 61003, Cedex 4, Montpellier 34184, France, \(^2\)Inserm U 1051, Institut des Neurosciences de Montpellier, Hôpital Saint Eloi, Montpellier 34091, France, \(^3\)Université de Montpellier, Montpellier 34091, France, \(^4\)Department of Molecular Pathology, The Centre for Cancer Biology, SA Pathology and Box 14 Rundle Mall Post Office, Adelaide, South Australia 5000, Australia, \(^5\)The Schools of Medicine and Molecular and Biomedical Science, The University of Adelaide, Adelaide, South Australia 5005, Australia and \(^6\)Department of Genetic Medicine and Development, University of Geneva Medical School and University Hospitals of Geneva, 4 Geneva 1211, Switzerland

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Before acquiring their mature state, cochlear hair cells undergo a series of changes in expression of ion channels. How this complex mechanism is achieved is not fully understood. Tmprss3, a type II serine protease expressed in hair cells, is required for their proper functioning at the onset of hearing. To unravel the role of Tmprss3 in the acquisition of mature K\(^+\) currents, we compared their function by patch-clamp technique in wild-type Tmprss3\(^{WT}\) and Tmprss3\(^{Y260X}\)-mutant mice. Interestingly, only outward K\(^+\) currents were altered in Tmprss3\(^{Y260X}\)-mutant mice. To determine by which mechanism this occurred, we compared the protein network of Tmprss3\(^{WT}\) and Tmprss3\(^{Y260X}\)-mutant mice using proteomic analysis. This led to the identification of a pathway related to potassium Kcnma1 channels. This pathway was validated by immunohistochemistry, focusing on the most downregulated protein that was identified as a cochlear Kcnma1-associated protein, APOA1. Finally, we show that, in contrast to Tmprss3\(^{WT}\), Kcnma1 channels were absent at the neck of inner hair cells (IHCs) in Tmprss3\(^{Y260X}\)-mutant mice. In conclusion, our data suggest that lack of Tmprss3 leads to a decrease in Kcnma1 potassium channels expression in (IHCs).

**INTRODUCTION**

The perception of sound relies on the proper functioning of the cochlea, the peripheral auditory organ. In mammalian cochlea, inner hair cells (IHCs) translate sound stimulation into graded receptor potentials. At the onset of hearing (P10-P12 in mice), IHCs undergo massive changes in their electrical properties (1,2), and notably, the maturation of hair cells correlates with the expression of large outward-rectifying potassium currents, \(I_{K,f}\), carried by Kcnma1 channels (3). The fast activation kinetics of these channels shape the time course and amplitude of the hair cell receptor potential (4), indicating that Kcnma1 channels in hair cells are critical for high-fidelity sound encoding. At the same time \(I_{K,n}\) current is expressed to set the resting potential of IHCs (5,6). This current is generated by KCNQ4 whose mutations are responsible for DFNA2 human deafness (7). Finally, a classical delayed-rectifier K\(^+\) current, name \(I_{K,s}\) shows slow activation kinetics and represents the slow component of the K\(^+\) outward current (3).

Mutations in the TMPRSS3 gene were shown to cause human autosomal recessive non-syndromic deafness (DFNB8/10) (8), characterized by bilateral, severe to profound hearing loss. To address the role of the protein in the cochlear physiology, we generated an ethynitrosurea (ENU) C3HeB/FeJ-mutant mouse (9). A T to A substitution in exon 7, resulting in a nonsense mutation at tyrosine 260 was identified (Tmprss3\(^{Y260X}\)) that would result in the production of a 194 amino acid truncated protein, deleted of most of its protease domain (9).

\(^\dagger\)These authors contributed equally to this work.

\(^*\)To whom correspondence should be addressed at: Inserm U1051, 80 rue Augustin Fliche, Montpellier 34091, France. Tel: +33 499636061; Fax: +33 499636020; Email: benjamin.delprat@inserm.fr

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The analysis of the Tmprss3Y260X-mutant mice revealed that Tmprss3 is essential for mouse cochlear hair cell survival at the onset of hearing and that Tmprss3 mutations induce deafness in mice (9) and in humans. Indeed, in the Tmprss3Y260X-mutant mice, hair cells degenerate drastically in 2 days (between P12 and P14). Thus, Tmprss3 is expressed in cochlear hair cells in the developing cochlea and at the onset of hearing, suggesting that Tmprss3 may be a hair cell-autonomous survival factor (9). We also identified a novel, longer Tmprss3 isoform with a 22 amino acid extension at the N-terminal, namely Tmprss3f. This isoform is expressed predominantly in the cochlea in contrast to the shorter isoform that is expressed broadly, suggesting a specific role of Tmprss3f in hair cell functioning and/or survival (9). Finally, the only known substrate of Tmprss3 that was observed in Xenopus oocytes (10) is the epithelial amiloride-sensitive sodium channel (ENaC).

Because IHC maturation is dependent on a change in ionic conductance expression, and given that Tmprss3 is known to activate ENaC channels in Xenopus oocytes, we would like to determine whether Tmprss3 was able to modulate the function of channels or channel regulatory proteins in IHCs. Using an integrated approach combining the patch-clamp technique, two-dimensional (2D) gels followed by mass spectrometry analysis, bioinformatic functional studies and immunohistochemistry, we showed, for the first time, that Tmprss3 loss of function leads to a decrease in the IHCs’ Kcnma1 channel plasma membrane expression. These alterations were associated with the downregulation of ApoA1 in IHCs, which is a protein identified as a Kcnma1 channel interacting partner expressed in IHCs (11).

Our data highlight a novel link between Kcnma1 channels and a serine protease. This unique finding suggests that this original relationship seems highly important for the normal physiology of IHCs and it should be significant for other essential body functions, where Kcnma1 channels play a key role, such as neurotransmission (12), blood flow (13), urine physiology of IHCs and it should be significant for other essential body functions, where Kcnma1 channels play a key role, such as neurotransmission (12), blood flow (13), urine flow (14) and immunity (15).

**RESULTS**

**Lack of fast K⁺ conductance in Tmprss3Y260X mouse IHCs**

Because ENaC, the only known substrate of Tmprss3, is not expressed in cochlear hair cells (16–18), whole-cell patch-clamp recordings were used to probe the K⁺ current in P13-old apical IHCs before degenerative changes occurred in this region of the cochlea (9). When depolarized, IHCs from Tmprss3WT showed an outward K⁺ current with a fast activating outward component, reminiscent of Kcnma1 currents (3,4,19) and a more slowly activating component (Fig. 1A). IHCs from Tmprss3Y260X showed a drastic reduction in the amplitude of the outward currents when measured 1 ms after the start of the depolarizing pulses (mostly recruiting Kcnma1 channels activation; 4.6 ± 0.4 nA at 30 and 2.27 ± 0.15 nA at 37 mV in Tmprss3WT and Tmprss3Y260X, respectively). In contrast, the amplitudes of the currents at the end of the depolarizing pulses were comparable between the genotypes (Fig. 1D). The difference of K⁺ outward current observed between the Tmprss3WT and Tmprss3Y260X could be explained by a delayed maturation. However, exocytosis triggered by Ca²⁺ currents was similar between Tmprss3WT and Tmprss3Y260X in P9-P11-old IHCs (Data not shown), suggesting normal hair cell development (20). Current-clamp recordings (Iinj = 0 pA) indicated no difference in the resting membrane potential of the IHCs between both genotypes (−69.4 ± 1.3 mV and −67.9 ± 2.1 mV for Tmprss3WT and Tmprss3Y260X IHCs, respectively).

**Differential proteomics**

To identify proteins that could regulate the expression of IHC K⁺ channels and that are linked to Tmprss3, P10 mouse cochlear extracts from Tmprss3WT and Tmprss3Y260X were analyzed by a proteomic approach using 2D-gel electrophoresis (2D-GE). We chose P10 because it represents a good compromise between the presumed absence of the apoptosis-related proteins that may be present in cochlear extracts at P12 due to the beginning of hair cell degeneration, and the presence of mature K⁺ currents that begin to appear at P10. After image analysis, more than 2000 spots were detected from cochlear extracts with a wide range of molecular weights (15–150 kDa) and isoelectric points (3–10 pI). To compare the Tmprss3WT and Tmprss3Y260X proteomes, it was
important to firstly assess the consistency of our dataset after 2D-GE separation. To this end, we used a dispersion tree approach (20), and the 10 2D-GE experiments were dispersed homogeneously on the 10 branches of the tree, which formed a circle (Supplementary Material, Fig. S1A). The homogeneity of the data was also confirmed by three other statistical methods: the Mahalanobis distance, principal component analysis and Pearson correlation (Supplementary Material, Fig. S1B–D).

In the comparative analysis of all protein spots between Tmprss3WT and Tmprss3Y260X, 77 distinct protein spots were differentially expressed (Fig. 2A). To visualize these protein spots, we used a heat map and HAC analysis using the JMP v8 software. Each column represents the data from one 2D-GE experiment. Rows represent individual spots. Raw data were mean centered, and the graduated scale color codes from green (low level of intensity) to red (high level of intensity).

Figure 2. Proteomic analysis and altered pathway in Tmprss3Y260X-mutant mice. (A) 2D-GE of proteins extracted from Tmprss3WT cochleae (left gel) and Tmprss3Y260X cochleae (right gel). Seventy-seven protein spots were differentially expressed, and protein spots were up or downregulated (example of spot regulation in the side inset). The proteins of interest identified by LC-MS/MS are shown on the gels by a circle. (B) Heat map and HAC of the 77 protein spots present in all cochlea 2D-GE. The intensity levels of the protein spots were visualized by a heat map and HAC analysis using the JMP v8 software. Each column represents the data from one 2D-GE experiment. Rows represent individual spots. Raw data were mean centered, and the graduated scale color codes from green (low level of intensity) to red (high level of intensity). (C) Network of identified proteins connected to Kcnma1. Nodes and edges are represented by gray circles and black lines, respectively. Identified proteins differentially expressed between Tmprss3WT and Tmprss3Y260X were highlighted using a graduated scale from green (downregulated) to pink (upregulated) according to their fold ratio. Proteins that were both up and downregulated in different protein spots were represented by a ratio of 1 (yellow). Some edges may represent a set of different PPIs found in the database. View of the merged network imported from Cytoscape using the IntAct and BioGRID databases.
spots further, we clustered the high-confidence dataset of 77 proteins spots. The dataset was then subjected to hierarchical clustering analysis generating the heat map and the dendogram shown in Figure 2B. Two main clusters of protein spots were clearly detected and differentially regulated. These two clusters corresponded to upregulated proteins in the presence of Tmprss3Y260X (high intensity shown in red) and to downregulated proteins (low intensity shown in green). Thirty-three protein spots were downregulated and 44 were upregulated in Tmprss3Y260X mice. All the differentially regulated protein spots were identified by liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS) analysis, and the results are summarized in the Supplementary Material, Table S1. One hundred and nineteen independent proteins were identified (after excluding redundant proteins) because of the presence of several spots containing more than one type of identification. Indeed, some proteins were present in full size form and/or in fragment form, such as vimentin and tubulin beta-5, and others were present in different pI forms such as transaldolase 1 and sparc. Moreover, proteins that were upregulated in a spot and downregulated in another one were represented by a fold ratio of one. Less than 1% of proteins identified in our screen are related to apoptosis, suggesting that the altered expression of the proteins is due to a lack of function of Tmprss3, rather than a degenerative process initiated by the IHCs.

Perturbed network identification

To evaluate the potential functional relationships between the set of identified proteins, we carried out a gene ontology (GO) enrichment analysis. We then used the Ontologizer software to perform a statistical analysis of the mapping of the 119 identified proteins over the functional GO terms (Supplementary Material, Table S2). Subsequently, we built a wider network by looking at connected proteins (first-degree neighbors) with our set of 119 identified proteins using the IntAct and BioGRID merge network. The average degree of connectivity with our set of 119 identified proteins using the IntAct and BioGRID databases. From the main subnetworks (160 proteins), only 4 highly connected ‘hub’ proteins were identified. These ‘hubs’ were Ywhab, Ywhaz, Dlh4 and Kcnma1 corresponding to the following proteins: 14-3-3 protein beta/alpha (Ywhab), 14-3-3 protein zeta/delta (Ywhaz), disks large homolog 4 (Dlg4/PSD95) and calcium-activated potassium channel sub-unit alpha-1 (Kcnma1) connecting 17, 5, 13 and 22 identified proteins, respectively (Fig. 2C and Supplementary Material, Fig. S2). Interestingly, Kcnma1 protein connected the largest number of identified proteins from comparative 2D-GE analysis between Tmprss3WT and Tmprss3Y260X.

APOA1 is downregulated in Tmprss3Y260X-mutant mice

Among the proteins identified by mass spectrophotometry, APOA1 was the most known cochlear interacting protein of Kcnma1, which was downregulated. To validate the APOA1 regulation by Tmprss3, immunohistochemistry was implemented on P13 cochlear explants. Using a polyclonal antibody, APOA1 (red) seemed to be expressed mainly in the IHCs of Tmprss3WT mice labeled with a monoclonal parvalbumin antibody (green) (Fig. 3A). This expression pattern is expected as APOA1 has been shown to be expressed in the cytoplasm of mouse IHC (11). In contrast, APOA1 labeling was fainter in the remaining Tmprss3Y260X IHCs (Fig. 3B). Indeed, the total intensity of the APOA1 signal inside each IHC is greater in Tmprss3WT mice than in Tmprss3Y260X mice (Fig. 3C). Because ApoA1 is known to interact with Kcnma1 channels (11), the downregulation of ApoA1 in Tmprss3Y260X mice suggests that Tmprss3 impairs the Kcnma1 channel’s interacome.

Kcnma1 channels are not clustered in puncta in Tmprss3Y260X mouse IHCs

It is well known that Kcnma1 channel immunoreactivity is clustered in puncta in the neck of mammalian IHCs (19). Therefore, it is possible to determine whether Kcnma1 channels are correctly expressed in the IHCs of Tmprss3Y260X-mutant mouse. To address this question, immunohistochemical staining of the Kcnma1 channel using a monoclonal anti-Kcnma1 antibody was used in the developing Tmprss3WT and Tmprss3Y260X organ of Corti in the same cochlear turn from P12 to P14 (Fig. 4). Indeed, as there were no more hair cells from P15 in Tmprss3Y260X-mutant mice, we did not pursue our study further after P14. At P12, the period corresponding to the onset of hearing in mice, some Tmprss3WT IHCs expressed Kcnma1 channels in their neck (Fig. 4A). In contrast, no Tmprss3Y260X IHCs expressed Kcnma1 channels at this stage (Fig. 4B). At P13, more Tmprss3WT IHCs expressed Kcnma1 channels at this stage (Fig. 4B). At P13, more Tmprss3WT IHCs expressed Kcnma1 channels (Fig. 4D), and signs of IHCs that had degenerated were observed. Finally, at P14, all Tmprss3WT IHCs expressed Kcnma1 channels, and the number of puncta increased per hair cell (Fig. 4C). As at P12, no P13 Tmprss3Y260X IHCs expressed Kcnma1 channels (Fig. 4D), but signs of IHCs that had degenerated were observed. Finally, at P14, all Tmprss3WT IHCs expressed Kcnma1 channels, and the number of puncta increased (Fig. 4E). At P14 in Tmprss3Y260X IHCs, the majority of IHCs had degenerated, but in the persisting IHCs, no Kcnma1 channel expression was observed (Fig. 4F). We quantified the number of IHCs expressing Kcnma1 channels in their neck depending on the cochlear turn and the age of the animals (see Fig. 4G). The number of IHCs expressing Kcnma1 channels is greater in the basal turn of the cochlea than in the apical one and increased from P12 to P14 as previously observed (19).

DISCUSSION

The lack of expression of Kcnma1 channels at the plasma membrane may be due to defects of various cellular processes.
One hypothesis may be that as the hair cells begin to degenerate, the consequence will be an altered expression of Kcnma1 channels. Our data show that only Kcnma1 channels are lacking—the delayed rectifier $K^+$ channel is still present as is the calcium channel Cav1.3 (this study). However, specific abolition of Kcnma1 channels when cochlear hair cells are degenerating has never been reported to our knowledge. Moreover, in Beethoven mutant mice, hair cells still have residual $I_{K,f}$ current just before their degeneration (21). Hence, it is unlikely that the specific lack of Kcnma1 channels is due to the degeneration of cochlear hair cells. Another hypothesis is that Tmprss3$^{Y260X}$ hair cell maturation fails or is delayed as IHCs from Tmprss3$^{Y260X}$ are smaller than WT littermate. Indeed, genetic ablation of the calcium channel Cav1.3 sub-unit (19), and its associated β-2 sub-unit (22), ER transmembrane protein Tmc1 (21), molecular motor Myo6 (23), actin-binding protein (24), or thyroid hormone receptor (25) leads to the absence of $I_{K,f}$ current because of a failure or a delay in cochlear hair cell maturation. However, transmission electron microscopy observations of Tmprss3$^{Y260X}$ IHCs show normal development until their degeneration [(presence of small numbers of ribbons/IHCs, mature synaptic contacts with only one ribbon/synapse, efferent synaptic contact onto lateral afferent under IHC and direct efferent synaptic contact onto OHC (9)], and normal $I_{ca}$ current amplitude and normal exocytosis were measured in Tmprss3$^{Y260X}$ when compared with Tmprss3$^{WT}$ (this study). All these observations suggest involvement of other mechanisms besides maturation failure or delay.

Observing our differentially expressed protein data, we identified proteins associated with Kcnma1 channels that may modulate the membrane expression of channels and transporters. Indeed, 15 out of 119 and 13 out of 119 of the proteins identified in the present screen were identified previously as Kcnma1 channel interacting partners that originated from membrane/cytoskeletal and cytoplasmic fraction, respectively (26,27). These data suggest that Tmprss3 is a part of Kcnma1 channel pathway and that loss of function of Tmprss3 interferes with proper Kcnma1 channel membrane expression. Interestingly, three proteins identified in our screen induce deafness either in humans (γ actin and serpinb6) or mice (β actin). ACTG1 (γ actin) was identified as the causative gene of DFNA20/26 human deafness (28) and in the syndromic deafness form of Baraitser-Winter Syndrome 2 (29). All DFNB20/26 patients display progressive, bilateral, sensorineural hearing loss that begins in the high frequencies. As age increases, the degree of hearing loss increases, with threshold shifts seen at all frequencies (28). In contrast, congenital or late-onset progressive hearing loss is a common feature of Baraitser-Winter syndrome 2. Actg1 is essential for the stereocilia maintenance in mice (30), but not their normal development. However, Tmprss3$^{Y260X}$-mutant mice have normal stereocilia (9). Interestingly, downregulation of Actg1, which interacts with Kcnma1, decreases Kcnma1 channel membrane expression in transfected CHO cells (27). As Actg1 is downregulated in our screen, it is tempting to speculate that Tmprss3 affects the distribution of Actg1 that in turn affects the proper expression of Kcnma1 channel at the inner hair cell plasma membrane. ACTB (β actin) mutations cause Baraitser-Winter Syndrome 2 (29). Actb is also required for stereocilia maintenance, and mutations in mice lead to progressive deafness (30). Like Actg1, Actb interacts with Kcnma1, and its downregulation in podocytes decreases Kcnma1 surface expression (31). As Actb is downregulated in our proteomic screen, this may lead to a decreased expression of Kcnma1 at the inner hair cell surface. Interestingly, Eps8, a protein with actin binding, bundling, and barbed-end capping activities is a novel component of the hair bundle that is localized predominantly at the tip of the stereocilia and is essential for its normal elongation and function (24). Surprisingly, IHCs from Eps8 knockout mice do not develop adult-type ion channels (lack of $I_{K,f}$ conductance linked to

Figure 3. APOA1 is downregulated in IHCs Tmprss3$^{Y260X}$-mutant mice. Cochlear explants were stained with anti-APOA1 antibody in P13 mice. (A) APOA1 (red) is expressed mainly in IHCs, identified by parvalbumin labeling (green) in Tmprss3$^{WT}$ cochlea. (B) Less staining could be observed in the remaining IHCs of Tmprss3$^{Y260X}$-mutant mouse. (C) Quantification of the intensity of APOA1 labeling in hair cells in Tmprss3$^{WT}$ (24%) and Tmprss3$^{Y260X}$-mutant mouse (7.6%). The percentage corresponds to the percentage of red color ($n = 3$ for Tmprss3$^{WT}$, $n = 4$ for Tmprss3$^{Y260X}$). The difference is statistically significant (two sample Wilcoxon test, $p < 0.05$).
Kcnma1 channels) as observed in Tmprss3Y260X mice, but $I_{\text{Ca}}$
current and exocytosis fail to mature in Eps8 knockout mice in
contrast to Tmprss3Y260X mice. These data suggest that the
same phenotype (lack of Kcnma1) is due to two different
mechanisms. Finally, SERPINB6-truncating mutation is asso-
ciated with DFNB91 deafness (32). Serpinb6 is a proteinase
inhibitor of cathepsin G and kallikreins. It has been shown
that Serpinb6 exhibits a prosurvival function in Caenorhabdi-
tis elegans (33). In developing and in adult cochlea, Serpinb6
is expressed in cochlear hair cells, suggesting a critical role of
the protein in hair cell physiology (32). Nevertheless, Serpinb6
has not yet been identified as a partner of Kcnma1, and no in-
formation could be found showing a link between Serpinb6
and surface membrane expression of Kcnma1.
The analysis of the proteomic approach underlies 14.3.3 β and 14.3.3 ζ as potential proteins functionally related to the Tmprss3Y260X mutation. Considering the IHC Kcnma1 channel plasma membrane alteration, it is tempting to speculate that 14.3.3 proteins may be involved. Indeed, it has been shown that 14.3.3 β, 14.3.3 ε, 14.3.3 η, 14.3.3 γ, 14.3.3 τ and 14.3.3 ζ interact with Kcnma1 channels and that downregulation of 14.3.3 γ induces an increase in Kcnma1 channel membrane expression in CHO cells (27). 14.3.3 β knockdown increases ENaC membrane expression (34), 14.3.3 β mediates forward trafficking to the plasma membrane of KCNK3 (35) and 14.3.3 ζ mediates trafficking to the plasma membrane of KA2 kainate receptors (36). Moreover, 14-3-3 proteins promote the cell surface transport of correctly assembled complexes (37). Interestingly, cleavage of 14-3-3 proteins, which is executed by at least two different mechanisms, probably involves caspase-3 and a still unknown protease beyond the caspase family in an isoform-specific manner (38) that could be Tmprss3. Of course, we cannot rule out the possibility that Tmprss3 directly controls the membrane expression level of Kcnma1 channels, as Tmprss3 and Kcnma1 channels co-localize in the ER of transfected cells.

This novel link between Kcnma1 channels and a type II serine protease may be highly significant in other organs, as Kcnma1 channels are widely expressed in neuronal and non-neuronal tissues, including epithelia, smooth muscle cells and sensory cells, where they couple membrane potential and intracellular Ca^{2+} concentration (39–41). Kcnma1 channels contribute to the repolarization of action potentials (APs) (42), mediate the fast phase after hyperpolarization following an AP (43–45), shape the dendritic Ca^{2+} spikes (46) and influence the release of neurotransmitters (47). Achieving a better understanding of the role of Kcnma1 channels is important not only for furthering our knowledge of the involvement of these channels in physiologic processes but also for pathophysiologic conditions that have been demonstrated by recent discoveries implicating these channels in neurologic disorders. Indeed, autism and mental retardation have been considered as an important factor for the proper functioning of hair cells.

MATERIALS AND METHODS
Animal handling
The care and use of animals followed the animal welfare guidelines of the ‘Institut National de la Santé et de la Recherche Medicale’ (Inserm), under the approval of the French ‘Ministère de l’Alimentation, de l’Agriculture et de la Pêche.’ All efforts were made to minimize the number of animals used and their suffering. Tmprss3Y260X mice were described previously by Fasquelle et al. (9).

Preparation of mouse inner ear extracts for proteomics analysis
Cochleae were isolated from 15 10-day-old mice of each genotype (WT and mutant) in cold PBS, and the bone was removed. The dissected cochleae were rinsed in water and wiped with paper before being placed in a microcentrifuge tube and weighed. The cochleae pellets were then frozen in liquid nitrogen and stored at -80°C. According to their weight, the pellets were resuspended in a defined volume of buffer (100 μL/10 mg of tissue) containing 8 M urea, 2 M thiourea, 4% w/v CHAPS, 65 mM dithioerythritol (DTE), 40 mM Tris and protease inhibitors (Roche Diagnostics, Meylan, France) and sonicated for 3 × 10 s pulses at 20 kHz (Vibracell 72446, Fisher Scientific, Illkirch, France). After solubilization at room temperature (RT) on a rotating wheel for 2 h, the samples were centrifuged at 20 000 g for 30 min, and the supernatants were collected and stored at -80°C until use. The amount of protein was estimated using a modified Bradford method (49).

2D-GE of proteins
Precast IPG strips (18 cm) with a non-linear immobilized pH 3–10 gradient were rehydrated with 70 μg of protein sample in 8 M urea, 2 M thiourea, 4% w/v CHAPS, 65 mM DTE, 0.0025% v/v bromophenol blue, and 1% v/v IPG buffer (3–10) overnight (50). Isoelectric focusing (IEF) was carried out on the Ettan™ IPGphor™ IEF system at 20°C using a gradient mode to a total amount of 50 kVh. After the first dimensional run, the proteins were reduced (65 mM DTT in 6 M urea, 50 mM Tris–HCl, pH 8.8, 30% v/v glycerol, 2% w/v SDS and 0.001 v/v bromophenol blue) and then alkylated for 10 min in a similar buffer containing 135 mM iodoacetamide instead of DTT. Subsequently, the proteins were separated in the second dimension on homemade 12% SDS-polyacrylamide gels using an ISO-DALT electrophoresis unit at a constant voltage of 120 V at 10°C overnight. Analytical gels were stained with a Sypro Ruby fluorescence dye (Bio-Rad, Hercules, CA, USA) according to the manufacturer’s instructions. At least five biologic replicates per genotype (Tmprss3WT and Tmprss3Y260X) were performed simultaneously to guarantee reproducible results. Each biologic sample was composed of a pool of six individual cochleae corresponding to three animals.

Image analysis
Gels were digitalized individually with a Typhoon 9400 laser scanner (GE Healthcare Life Sciences, Piscataway, NJ, USA) at 50 μm resolution with a photo multiplier tube voltage adjusted for maximum range without signal saturation. Images were stored as 16-bit gel files and then analyzed using the Progenesis Samespot® software v3.0 (Nonlinear Dynamics, Durham, UK). Images were warped for accurate alignment and spots were then detected automatically. Progenesis
Samespot® is based on the concept of recursive gel matching, which means that each gel in a matching set is used recursively as ‘reference gel’ once during the matching process. The quality of the automatic match was evaluated critically in each case, and if necessary, corrections were made manually.

Statistics
All statistics and figures were computed using the 'R/Bioconductor' statistical open source software (51) and the JMP v8 software (SAS Institute, Inc.). The differential intensity levels of protein spots between the Tmprss3WT and Tmprss3Y260X samples were analyzed by the non-parametric Wilcoxon rank sum test (also called the Mann–Whitney U-test). A P-value less than 0.05 was considered statistically significant. The area under the curve (AUC) receiver operating characteristic (ROC) was also calculated with the ROC package, and an AUC ROC value greater than 0.800 was considered significant.

Hierarchical ascendant clustering (HAC) analysis
Hierarchical ascendant clustering (HAC) is a method of cluster analysis based on a pairwise distance matrix, which builds a hierarchy of clusters with sequentially agglomerative and divisive approaches. We used this method to organize the map and to group the spots according to the nearest level of intensity. For this analysis, raw data were mean centered, and the Pearson correlation matrix and average linkage were chosen as parameters.

Protein identification
Spots of interest were excised from preparative gels with a Propic robot (Perkin-Elmer, Wellesley, MA, USA). All subsequent steps were done automatically using a Multiprobe II robot (Perkin-Elmer). Spots were first washed with 300 μl of 2% formic acid. After pre-concentration, they were then dehydrated twice using 300 μl of 100% CH3CN and finally dried at 37°C for 1 h. Eight microliters of a trypsin solution (Sequencing grade modified trypsin, Promega, Madison, USA) at a concentration of 0.0125 μg/μl in 25 mM NH4HCO3 was added to every spot. Digestion was performed overnight at 37°C and stopped by the addition of 2 μl of 2% formic acid. Digests were sonicated in an ultrasonic bath for 10 min, and supernatants were transferred into HPLC polypropylene tubes.

Protein digests were analyzed using a high-capacity ion trap mass spectrometer (Esquire HCT; Bruker Daltonik GmbH, Bremen, Germany) and interfaced with a nano-HPLC Chip-Cube system (Agilent Technologies, Santa Clara, USA). The chips contained both the pre-column and the column (Zorbax 300SB-C18; Agilent Technologies). Samples were first loaded onto the 4 mm enrichment cartridge at a flow rate of 0.3 μl/min using a 15 min linear gradient from 3 to 80% acetonitrile in 0.1% formic acid and eluted into the mass spectrometer. A capillary voltage of 1.8–2.1 kV in the positive ion mode was used together with a dry gas flow rate of 4.5 l/min at 250°C. A first full-scan mass spectrum was measured in the 310–1800 m/z range, followed by a second scan at higher resolution to measure precisely the mass of the three major ions in the previous scan. Finally, a third scan was performed to acquire the collision-induced MS/MS spectra of the selected ions. MS/MS raw data were analyzed using the data analysis software (Bruker Daltonik GmbH, Bremen, Germany) to generate the peak lists. The NCBI non-redundant database (NCBIrr, release 20101018) was queried locally using the Mascot search engine (v. 2.2.04; Matrix Science, London, UK) with the following parameters: Mus for the taxonomy, trypsin as the enzyme, one missed cleavage allowed, carbamidomethylation of Cysteine as a fixed modification, oxidation of Methionine as a variable modification, and 0.6 Da mass accuracy in both MS and MS/MS. Under these conditions, individual ion scores above 38 indicated identity or extensive homology (P < 0.05), and proteins were validated once they showed at least 1 peptide over this threshold.

Protein interaction data and network visualization
The gene names of the identified proteins were used to perform GO enrichment analysis using the Ontologizer software. This Java Web Start application allows the statistical analysis of sets of proteins or genes of interest mapped onto GO terms (http://compbio.charite.de/ontologizer) (52). GO annotation files (gene_association.wb and gene_ontology_dictionary.obo) were downloaded from www.geneontology.org in October 2011. The ontologizer was run using the following parameters: biologic process, parent–child union and the Benjamin–Hochberg statistical test. Biologic processes showing P-values below 0.05 were considered statistically significant. The ontologizer allows visualization of data as a directed acyclic graph highlighting all significantly enriched GO terms, in turn linked to the genes/proteins of interest.

PPI data were combined with our set of identified proteins to build a functional network. Full PPI datasets were available from two different open source literature-curated protein interaction databases: IntAct (http://www.ebi.ac.uk/intact/index.jsp; dated March 28 2011) and the Biological General Repository for Interaction Datasets (BioGRID; http://thebiogrid.org/; version 3.1.74). PPI networks were generated using Cytoscape (53), an open source software (http://www.cytoscape.org/; version 2.8.1). The gene names of all identified proteins were imported into IntAct via its Cytoscape plug-in. BioGRID datasets from Mus musculus (file BIOGRID-ORGANISM-Mus_musculus-3.1.74.tab) were imported into Cytoscape via the BioGRID plug-in (BioGridPlugin2). Proteins were represented by nodes and the interactions by edges. Then, for each of identified proteins, the direct interacting neighbors were identified (length = 1). Finally, the PPI networks from the IntAct and BioGRID databases were merged with an advanced merge network plug-in and displayed with Cytoscape using the edge-unweighted spring embedded style. The selection of subnetworks was based on the degree of connectivity of a significant number of identified proteins. Highly connected proteins were named ‘hubs proteins.’
Immunohistochemistry and confocal imaging

Cochleae were extracted from the temporal bones of 12-, 13- or 14-day-old mice and fixed in cold 4% paraformaldehyde for 1 h with shaking. After several washes in PBS, organs of Corti were isolated from the stria vasularis and modiolus and incubated in a goat serum dilution buffer (GSDB) (16% normal goat serum, 450 mM NaCl, 0.3% Triton X-100, and 20 mM phosphate buffer, pH 7.4) at RT for 1 h. Anti-Kcnma1 rabbit antibody (Sigma Aldrich, St. Louis, MO, USA), anti-APOA1 goat antibody (Novus Biologicals, Cambridge, UK) that was successfully used in mouse (54,55) and anti-parvalbumin mouse antibody (Swant, Marly, Switzerland) were diluted in GSDB at 1/2000, 1/500 and 1/750, respectively and applied overnight at 4°C with shaking. After washing with washing buffer (450 mM NaCl, 20 mM phosphate buffer pH 7.4 and 0.3% Triton X-100) (6x 10 min with shaking), the organs of Corti were incubated with goat anti-rabbit and goat anti-mouse antibodies, respectively, coupled with Alexa Fluor-594 and 488 (Jackson ImmunoResearch Laboratories, Suffolk, UK) diluted at 1/3000 in GSDB for 1 h 30 at RT with shaking and protection from light. A 5 min supplementary incubation was carried out after the addition of Hoechst (FluoProbes, Interchim, Montluçon, France) at 1/200. After 10 min washes each in a washing buffer, the organs of Corti were washed for 10 min in 5 mM phosphate buffer, pH 7.4, placed onto a glass microscope slide in a DaKo fluorescent mounting medium (DakoCytomation, Trappes, France) and covered with a thin glass coverslip. Confocal images were acquired using a laser scanning confocal microscope LSM 510 (Zeiss, LE PECQ, France) with argon and helium–neon lasers for excitation. Oil immersion objectives with a magnification of 63× were used, and Z-stacks were acquired using a step size of 1.5 μm. Three-dimensional reconstruction images were made using the Imaris software. No immunoreactivity was detected in the absence of the primary antibodies.

The intensity of APOA1 immunolabeling was quantified from red green blue confocal images using custom-made program written in MATLAB (The Mathworks). The APOA1-labeled voxels (red channel) localized in IHC cytoplasm (green channel) were (i) isolated using parvalbumin immunolabeling as a mask and (ii) used to calculate the mean intensity. Mean intensities were calculated in Tmprss3WT (n = 3) and Tmprss3215305 animals (n = 4) and compared with Mann–Whitney–Wilcoxon test. These numbers correspond to the number of measurements. Each measurement contains at least five hair cells.

As mouse cochlea length has been found to average 6 mm (56), including the basal (3 mm), the middle (2 mm) and the apical (1 mm) coils, and as the IHCs have already degenerated at P12, the counting of the number of IHCs expressing Kcnma1 channels was achieved in middle turn (separated into two, M1 and M2) and in the apical turn (A).

Patch-clamp recordings

After cervical dislocation of the Tmprss3WT and Tmprss3215305 mice (postnatal day 13, P13), IHCs of the apical coil of freshly dissected organs of Corti were patch clamped at RT in whole-cell configuration. The pipette solution contained (in mM) 135 KCl, 10 Hepes buffer, 1 MgCl2, 2 Mg-ATP, 0.3 Na-GTP and 5 ethyleneglycol-bis(β-aminoethyl ether)-N,N,N',N'-tetraacetic acid. The extracellular solution contained (in mM) 144 NaCl, 5.8 KCl, 0.9 MgCl2, 10 HEPES, 1.3 CaCl2 and 10 d-glucose. The solutions were adjusted to pH 7.2 with osmolarities between 290 and 310 mOsmol/l. All chemicals were obtained from Sigma (Sigma Aldrich). The EPC-10 amplifier (HEKA, Lambrecht, Germany), controlled by the Patchmaster software (HEKA), was used for recording measurements. For K+ currents, series resistance (Rseries) compensation was applied to 60%. The voltage was corrected offline for the voltage drop across uncompensated series resistance. Currents were low-pass filtered at 5 kHz and sampled at 50 kHz. All voltages were corrected for liquid junction potentials calculated between the pipette and the bath (4 mV). Currents were leak corrected using a p/10 protocol (usually 10 leak pulses with amplitudes of 20% of the original pulse from a holding potential of −104 mV). Cells that displayed a membrane current exceeding −50 pA at −74 mV were discarded from the analysis.

AUTHORS’ CONTRIBUTIONS

B.D. designed research; L.M., L.F., R.N. and N.S. performed research; L.M., L.F., R.N. and N.S. analyzed the data; M.G. and H.S.S. contributed new reagents; and L.M., R.N., F.M., M.G., H.S.S., J.L.P. and B.D. wrote this paper.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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