**Tmc1 Point Mutation Affects Ca^{2+} Sensitivity and Block by Dihydrostreptomycin of the Mechanoelectrical Transducer Current of Mouse Outer Hair Cells**

Laura F. Corns, Stuart L. Johnson, Corné J. Kros, and Walter Marcotti

1Department of Biomedical Science, University of Sheffield, Sheffield, South Yorkshire S10 2TN, United Kingdom, 2Sussex Neuroscience, School of Life Sciences, University of Sussex, Falmer, Brighton BN1 9QG, United Kingdom, and 3Department of Otorhinolaryngology, University Medical Center Groningen, University of Groningen, 9700 RB Groningen, The Netherlands

The transduction of sound into electrical signals depends on mechanically sensitive ion channels in the stereociliary bundle. The molecular composition of this mechanoelectrical transducer (MET) channel is not yet known. Transmembrane channel-like protein isoforms 1 (TMC1) and 2 (TMC2) have been proposed to form part of the MET channel, although their exact roles are still unclear. Using Beethoven (Tmc1Bth/Bth) mice, which have an M412K point mutation in TMC1 that adds a positive charge, we found that Ca^{2+} permeability and conductance of the MET channel of outer hair cells (OHCs) were reduced. Tmc1Bth/Bth OHCs were also less sensitive to block by the permeant MET channel blocker dihydrostreptomycin, whether applied extracellularly or intracellularly. These findings suggest that the amino acid that is mutated in Bth is situated at or near the negatively charged binding site for dihydrostreptomycin within the permeation pore of the channel. We also found that the Ca^{2+} dependence of the operating range of the MET channel was altered by the M412K mutation. Depolarization did not increase the resting open probability of the MET current of Tmc1Bth/Bth OHCs, whereas raising the intracellular concentration of the Ca^{2+} chelator BAPTA caused smaller increases in resting open probability in Bth mutant OHCs than in wild-type control cells. We propose that these observations can be explained by the reduced Ca^{2+} permeability of the mutated MET channel indirectly causing the Ca^{2+} sensor for adaptation, at or near the intracellular face of the MET channel, to become more sensitive to Ca^{2+} influx as a compensatory mechanism.

**Key words:** adaptation; calcium; development; hair cell; mechanotransduction; TMC1

**Introduction**

The sense of hearing relies on the transduction of sound stimuli into electrical signals, which is performed by mechanically gated ion channels located at the tips of the shorter stereocilia (Beurg et al., 2009). Stereocilia are specialized microvilli forming staircase-like structures, the hair bundles, that project from the upper surface of hair cells. The stereocilia within a hair bundle are...
interconnected by extracellular linkages, the most important of which for the purpose of mechanoelectrical transduction is the tip link (Pickles et al., 1984; Assad et al., 1991; Goodyear et al., 2005). Tip links connect the stereocilia in the direction of optimal mechanical sensitivity, enabling them to transmit force to the mechanoelectrical transducer (MET) channels during bundle displacement, thus modulating their open probability. Although the molecular identity of the MET channel is still uncertain, recent studies have identified two molecules (TMHS/LHFPL5 and TMIE) that are implicated in mechanoelectrical transduction and directly interact with protocadherin-15 (PCDH15), which connects the lower end of the tip link into the stereociliary top (Xiong et al., 2012; Zhao et al., 2014). However, neither of these proteins appears to be the pore-forming subunit of the MET channel (Xiong et al., 2012; Zhao et al., 2014). TMC1 and TMC2 are localized at the tips of the shorter stereocilia (Kurima et al., 2015), and are the site for mechanoelectrical transduction, and are candidate subunits of the MET channel (Kawashima et al., 2011; Pan et al., 2013). However, recent reports have provided some evidence for a more accessory subunit role for both TMC1 and TMC2, mainly based on the fact that a mechano-gated current could still be elicited in TMC1/TMC2 double knock-outs (Bourg et al., 2014). However, this undefined mechano-gated current can also be elicited in hair cells from normal mice in the absence of tip links and has biophysical and pharmacological properties that are somewhat distinct from those of the MET channel (Marcotti et al., 2014). If this anomalous mechano-gated channel is indeed the MET channel precursor in cochlear hair cells, then TMC1 and TMC2 could be required to functionally mature channel pore and/or vestibule facing the extracellular side of the channel.

The Beethoven (Bth) mutant mouse is a model for progressive hearing loss (DFNA36) caused by a single dominant missense mutation of a methionine to a lysine at position 412 of TMC1 (Kurima et al., 2002; Vreugde et al., 2002; Nakaniishi et al., 2014). In this study, we have used the Beethoven mouse and a double-mutant Beethoven/Tmc2 mouse model to provide additional evidence for a direct role of TMC1 and TMC2 in mechanoelectrical transduction in mouse outer hair cells (OHCs). Our results indicate that the M412K point mutation in Beethoven mice reduces the permeability of the MET channel for Ca$^{2+}$ and the permeant blocker dihydrostreptomycin (DHS). The Bth mutation also abolished the normally observed increase of the resting MET current on OHC depolarization. Increasing the extracellular concentration from its endolymphatic concentration of ~0.04 to 0.1 mM or decreasing intracellular Ca$^{2+}$ buffering caused a decrease in resting open probability of the MET current in Bth mutant OHCs but not in wild-type controls. We argue that these observations can be most parsimoniously explained by assuming that the reduced Ca$^{2+}$ permeability indirectly causes the Ca$^{2+}$ sensor for adaptation of the MET channel to become more sensitive to Ca$^{2+}$ influx.

**Materials and Methods**

**Animals and genotyping.** All experiments were performed in accordance with Home Office regulations under the Animals (Scientific Procedures) Act 1986 and following approval by the University of Sheffield Ethical Review Committee. The Beethoven (Tmc1Bth−/− and Tmc1Bth+/−) mice were maintained on a C57Bl6/FeJ background. Tmc1Bth−/− encodes a point mutation at residue 412 that is predicted to cause a methionine to lysine substitution (Vreugde et al., 2002). Tmc2 knock-out mice were obtained from The Jackson Laboratory (B6.129–Tmc2−/−H11001). Beethoven and Tmc2−/− mice were genotyped as described previously (Marcotti et al., 2006; Pan et al., 2013, respectively). The Beethoven mice were obtained from Karen Steel (Kings College, London, UK).

**Tissue preparation.** OHCs were studied in acutely dissected organs of Corti from postnatal day 2 (P2) to P12. Mice of either sex were killed by cervical dislocation, the cochlea removed, and the organ of Corti dissected in the extracellular solution composed of the following (in mM): 135 NaCl, 5.8 KCl, 1.3 CaCl$_2$, 0.9 MgCl$_2$, 0.7 NaH$_2$PO$_4$, 5.6 t-glycine, 10 HEPES-NaOH, and 2 Na-pyruvate. Amino acids and vitamins (Eagle’s MEM) were also added from concentrates (pH 7.5; osmolality, 308 mOsmol/kg). The dissected apical and basal coils of the organ of Corti were transferred to a microscope chamber containing extracellular solution and viewed through a long-working-distance 63× water-immersion objective on a Leica DMLS microscope.

**Whole-cell patch clamp.** MET currents were recorded at room temperature (20–25°C) using an Optopatch amplifier (Cairn Research). Patch pipettes, with resistances of 2–4 MΩ, were pulled from soda glass capillaries, and the shank of the electrode was coated with surf wax (Mr. Zoggs Sex Wax). For most experiments, except those measuring Ca$^{2+}$ selectivity, pipettes were filled with an intracellular solution composed of the following (in mM): 106 t-glutamic acid, 20 CsCl, 10 Na$_2$-phosphocreatine, 3 MgCl$_2$, 1 EGTa-CsOH, 5 NaATP, 5 HEPES, and 0.3 TGP (adjusted to pH 7.28 with 1 mM CsOH; osmolality, 294 mOsmol/kg). Data acquisition was performed using pClamp software (Molecular Devices) using a Digidata 1440A. Data were filtered at 5 kHz (eight-pole Bessel). Offline data analysis was performed using Origin software (OriginLab). Membrane potentials were corrected for a liquid junction potential of ~11 mV measured between electrode and bath solution.

**Hair bundle stimulation.** MET currents were elicited using a fluid jet from a pipette driven by a 25-mm diameter press electric disc (Kros et al., 1992; Corns et al., 2014). The fluid jet was preferred to a rigid probe (Peng et al., 2013) because it does not require any previous contact with the hair bundle, which could affect their resting position, and provides a more homogeneous displacement of stereocilia within a bundle compared with the stiff probe (Corns et al., 2014; Nam et al., 2015). The distance of the pipette tip from the bundle was adjusted to elicit a maximal MET current. Mechanical stimuli were applied as steps or saturating 50 Hz sinusoids (filtered at 1 kHz, eight-pole Bessel). Hair bundle motion during fluid jet stimulation was quantified by projecting an image of the OHC bundle onto a pair of photodiodes (LD 2–5; Centronics) at 360× magnification (Corns et al., 2014). This procedure is difficult to perform during MET recordings because it requires optimal orientation of the preparation to produce a bright bundle. Therefore, bundle displacement was not routinely calibrated and was converted as reported previously (10 nm/V; Corns et al., 2014) for the standard recording conditions: fluid jet pipette tip with a diameter of 8–10 μm positioned at ~8 μm from the bundles.

**Extracellular and intracellular application of the different solutions.** To test the effect of low Ca$^{2+}$ on the MET channel in vivo, the hair bundle was superfused with either low-millimolar Ca$^{2+}$ (0.1 mM), which is in the range of the predicted immature endolymphatic Ca$^{2+}$ (Johnson et al., 2012), or adult endolymph-like Ca$^{2+}$ (0.04 mM; Bosher and Warren, 1978; Salt et al., 1989). The solution containing 0.1 mM Ca$^{2+}$ was as follows (in mM): 147 NaCl, 5.8 KCl, 0.1 CaCl$_2$, 0.7 NaH$_2$PO$_4$, 2 Na-pyruvate, 5.6 $t$-glycine, and 10 K-HEPES-NaOH, pH 7.5. That with 0.04 mM Ca$^{2+}$ contained the following (in mM): 137 NaCl, 5.8 KCl, 0.04 CaCl$_2$ (buffered with N-(2-hydroxyethyl) ethylene-diamine triacetic acid: HEDTA), 5.6 $t$-glycine, and 10 Na-HEPES, pH 7.5. To investigate the effects of DHS (Sigma), 100 mM stock solutions of DHS (molecular weight, 730.7) were prepared in the requisite intracellular or extracellular solution. During the recordings, all test solutions were present in the fluid jet and also superfused via a pipette positioned orthogonally to the axis of mechanical sensitivity of the hair bundle, so the flow did not directly stimulate the stereocilia. For every extracellular solution change (i.e., changing from the standard 1.3 mM Ca$^{2+}$ to either 0.1 or 0.04 mM Ca$^{2+}$), the hair bundle was superfused with the new solution, and, before their stimulation, the fluid jet was filled with the same solution by suction through its tip using a negative gravity pressure system built in the fluid jet. Although this procedure allows to record the control MET current before the application of the different test solutions, the de-
gree of solution exchange in the fluid jet is unknown. To verify the reliability of our solution-exchange procedure in the fluid jet, we performed some experiments (21 of the 49 recordings used for Fig. 5) in which MET current was recorded with the fluid jet already prefilled with the different Ca2+ solutions. Because the two experimental procedures provided similar results, the data were pooled together in Figure 5.

To test whether MET current in mutant mice was regulated directly by the free Ca2+ inside the stereocilia, we investigated the influence of changing the intracellular Ca2+ buffering capacity. For these experiments, different concentrations (0.1–10 mM) of the fast Ca2+ buffer BAPTA (Invitrogen) was used instead of 1 mM EGTA (Fluka) in the above Cs-based intracellular solution. For the experiments investigating the effects of the drug were added to the Cs-glutamate intracellular solution. When the concentration of BAPTA or DHS used was >1 mM, Cs-glutamate was adjusted to keep the osmolality constant.

**Calcium selectivity.** The calcium selectivity of the MET channel was calculated by measuring the reversal potential of calcium (Vrev,Ca2+) using a CsCl-based intracellular solution [in mM: 135 CsCl, 3 MgATP, 10 Tris phosphocreatine, 1 EGTA-CaOH, 10 HEPES-CaOH, pH 7.2 (osmolality, 293 mOsmol/kg)] and a high Ca2+ extracellular solution (in mM: 100 CaCl2, 20 N-methylglucamine, 6 Tris, 10 r-glucose, adjusted to pH 7.4 with HCl), which was applied as described above. Reversal potentials were corrected for a liquid junction potential of −9 mV. The relative permeability, PCa/PNa, was calculated from the Goldman–Hodgkin–Katz equation:

\[
P_{\text{Ca}}/P_{\text{Na}} = \left[1 + \exp\left(V_{\text{Na}} - V_{\text{rev}} \right) / RT\right] / \left[1 + \exp\left(V_{\text{Ca}} - V_{\text{rev}} \right) / RT\right]
\]

where the binding constants, \(k_1\) and \(k_2\), rate constants are voltage dependent, and the Hill coefficient \(n_H = 1\). In brief, the voltage dependence of the block is expressed in four parameters: \(E_0\), the free energy of the drug binding site at 0 membrane potential; \(\delta_0\), the fractional electrical distance of the site across the membrane from the extracellular side; \(\Delta E = E_1 - E_0\), the difference between the intracellular and extracellular, respectively, free energy levels of the two barriers at 0 membrane potential; and \(\Delta \delta = \delta_1 - \delta_0\), the fractional electrical distance between the barriers. Fits using this model and assuming a Hill coefficient of 1 (Marcotti et al., 2005) yield values for \(\Delta E\), \(E_0\), \(\Delta \delta\), and \(\delta_0\), where the binding
Calcium selectivity of the MET channel is reduced in Tmc1Bth/Bth mice. A,B, MET currents recorded from P6 apical OHCs from Tmc1+/+ (A) and Tmc1Bth/Bth (B) mice in response to a 50 Hz sinusoidal force stimulus to the hair bundles superimposed to a voltage ramp (288 mV/s) from −129 to +91 mV nominal values (top panels). Arrows indicate the reversal potential of the MET current. C,D, Average MET current–voltage relationships recorded in apical (C) and basal (D) OHCs from Tmc1+/+ and Tmc1Bth/Bth mice in the voltage region around reversal potential. Data are approximated with a linear regression line, with the following slope: apical, 3.0 pA/mV in Tmc1+/+ and 2.1 pA/mV in Tmc1Bth/Bth; and basal, 3.4 pA/mV in Tmc1+/+ and 2.0 pA/mV in Tmc1Bth/Bth mice. Ca2+ reversal potential (left) and Ca2+ selectivity was significantly different (p < 0.0001) from that of Tmc1+/+ and Tmc1Bth/Bth cochleae is shown in Figure 2, C and D, respectively. The reversal potential of the MET current (Fig. 2E, left) was measured by fitting the I–V relationships from individual OHCs around the zero-current level with linear regression lines, which also allowed the calculation of the relative permeability, P_{Ca}/P_{Cs} (see equation in Materials and Methods; Fig. 2E, right). We found that the MET current reversal potential and relative Ca2+ permeability were similar between apical and basal OHCs, but both were reduced significantly (p < 0.0001) in Tmc1Bth/Bth OHCs compared with Tmc1+/+ OHCs (Fig. 2E). The smaller Ca2+ permeability in Tmc1Bth/Bth was also associated with a reduced MET current slope conductance over the entire voltage range investigated (by ~30% for apical and by ~40% for basal OHCs; Fig. 2C,D, respectively). In Tmc1Bth/+ heterozygotes, the Ca2+ selectivity was significantly different (p < 0.0001) from that of Tmc1+/+ and Tmc1Bth/Bth in both apical and basal OHCs (one-way ANOVA; post hoc test was p < 0.001 between Tmc1+/+ and Tmc1Bth/+ in both cochlear regions, and p < 0.01 and p < 0.05 between Tmc1Bth/+ and Tmc1Bth/Bth in apical and basal regions, respectively). Similar reductions in Ca2+ selectivity were also obtained elicited by displacing their hair bundles with a piezoelectric fluid jet stimulator (Kros et al., 1992, Corns et al., 2014). Saturating bundle displacement in the excitatory direction (i.e., toward the taller stereocilia) elicited large inward MET currents in OHCs from Tmc1+/+ and Tmc1Bth/Bth mice (Fig. 1A,B) and Tmc1Bth/+ mice (not data shown). Inhibitory hair bundle stimulation (negative driver voltage) shut off the small fraction of current flowing at rest (Fig. 1A,B, arrowheads). Comparably large MET currents were also elicited in OHCs from double-mutant mice in which Tmc2 was absent (Fig. 1C,D). The maximum amplitude of the MET current was similar at least up to P11 in all genotypes tested (Fig. 1E). These data demonstrate that the Beethoven mutation in Tmc1 and loss of Tmc2, which is only expressed up to P8–P10 in apical OHCs (Kawashima et al., 2011), do not impair the size of the MET currents.

The ionic selectivity of the MET channel in Beethoven mice was determined by measuring its selectivity for Ca2+ over other cations. The hair bundles of OHCs from Tmc1+/+ and Tmc1Bth/Bth mice were displaced using saturating stimuli superimposed on a voltage ramp (Fig. 2A,B) under conditions in which Ca2+ and Cs+ were the only permeant cations in the extracellular and intracellular solutions, respectively. The size of the MET currents in OHCs from both the apical and basal regions of the Tmc1+/+ and Tmc1Bth/Bth cochlea is shown in Figure 2, C and D, respectively. The reversal potential of the MET current (Fig. 2E, left) was measured by fitting the I–V relationships from individual OHCs around the zero-current level with linear regression lines, which also allowed the calculation of the relative permeability, P_{Ca}/P_{Cs} (see equation in Materials and Methods; Fig. 2E, right). We found that the MET current reversal potential and relative Ca2+ permeability were similar between apical and basal OHCs, but both were reduced significantly (p < 0.0001) in Tmc1Bth/Bth OHCs compared with Tmc1+/+ OHCs (Fig. 2E). The smaller Ca2+ permeability in Tmc1Bth/Bth was also associated with a reduced MET current slope conductance over the entire voltage range investigated (by ~30% for apical and by ~40% for basal OHCs; Fig. 2C,D, respectively). In Tmc1Bth/+ heterozygotes, the Ca2+ selectivity was significantly different (p < 0.0001) from that of Tmc1+/+ and Tmc1Bth/Bth in both apical and basal OHCs (one-way ANOVA; post hoc test was p < 0.001 between Tmc1+/+ and Tmc1Bth/+ in both cochlear regions, and p < 0.01 and p < 0.05 between Tmc1Bth/+ and Tmc1Bth/Bth in apical and basal regions, respectively). Similar reductions in Ca2+ selectivity were also obtained
in double-mutant mice in which Tmc2 was also absent (Fig. 2E, right). In both cases, the Ca\(^{2+}\) selectivity of the OHC MET channel was more than halved in Tmc1\(^{Bth/Bth}\) relative to wild type; a comparable reduction in Ca\(^{2+}\)/H\(^{11000}\) selectivity was reported in OHCs (Beurg et al., 2015) and inner hair cells (IHCs; Pan et al., 2013) of Bth mutants relative to wild type.

Effects of depolarization and extracellular Ca\(^{2+}\) on the MET current of OHCs from Beethoven mice

Calcium has been reported to modulate the fraction of the MET current activated in the absence of bundle stimulation. Although increasing Ca\(^{2+}\) influx through the MET channels promotes closing of some of the channels, measures to reduce Ca\(^{2+}\) influx into the channel, by either lowering its extracellular concentration or depolarizing the cell to near the Ca\(^{2+}\)/H\(^{11000}\) equilibrium potential, have been reported to increase channel open probability (Assad et al., 1989; Crawford et al., 1991; Ricci et al., 1998; Corns et al., 2014). Large MET currents could be recorded from OHCs of Tmc1\(^{+/+}\) (Fig. 3A) and Tmc1\(^{Bth/Bth}\) (Fig. 3B) mice at both negative (−121 mV) and positive (+99 mV) membrane potentials. OHC membrane depolarization caused the MET current to decrease in size at first and then reverse near 0 mV to become

**Figure 3.** Fraction of MET channels open at rest is reduced in Tmc1\(^{Bth/Bth}\) mice. A, B, Saturating MET currents in P9 apical OHCs from Tmc1\(^{+/+}\) (A) and Tmc1\(^{Bth/Bth}\) (B) mice in response to a 50 Hz sinusoidal force stimulus to the hair bundles at membrane potentials of −121 and +99 mV. The arrowheads and arrows indicate the closure of the transducer channels at −121 and +99 mV, respectively. \(V_{\text{Piezo}}\) indicates the driver voltage to the fluid jet. C, Peak-to-peak MET current–voltage curves obtained from five Tmc1\(^{+/+}\) and seven Tmc1\(^{Bth/Bth}\) OHCs (P9–P10) using 1.3 mM extracellular Ca\(^{2+}\). D, E, Resting open probability (\(P_\text{open}\)) of the MET channel at increasing membrane potentials from −121 to +99 mV in Tmc1\(^{+/+}\) (\(n = 6\)) and Tmc1\(^{Bth/Bth}\) (\(n = 8\)) mice. F, G, Resting \(P_\text{open}\) of the MET channel at the holding potentials from −121 (F) and +99 mV (G) from P2 to P12 from Tmc1\(^{+/+}\) and Tmc1\(^{Bth/Bth}\) mice. Number of OHCs tested at −121 mV (F) and +99 mV (G) are as follows: Tmc1\(^{+/+}\), 1, 3, 7, 6, 14, 3, 3, 3, 3; Tmc1\(^{Bth/Bth}\), 0, 0, 5, 4, 8, 5, 5, 3, 0.
outward at positive potentials (Fig. 3C), in accordance with the nonselective permeability of MET channels to cations (Ohmori, 1985). The fraction of the MET current open at rest is usually larger at positive compared with negative membrane potentials (Fig. 3A,D; Tmc1+/+). We found that Tmc1+Bth/Bth OHCs did not show any increase in the resting open probability (P_open) of the MET channel with membrane depolarization (Fig. 3B,D). Similar findings were also seen in the double-mutant mice in which Tmc2 was absent (Fig. 3E). At −121 mV, the resting P_open was relatively constant up to P12 in both Tmc1+/+ and Tmc1+Bth/Bth OHCs (Fig. 3F). At positive membrane potentials (+99 mV), although the resting P_open in Tmc1+/+ OHCs gradually increased with development (p < 0.0001, one-way ANOVA), that of Tmc1+Bth/Bth OHCs was found to be significantly smaller (p < 0.0005, two-way ANOVA) than that of Tmc1+/+ and relatively unchanged during the same time window (Fig. 3G). These findings suggest that the mutation in Beethoven mice strongly reduces the normal effect of depolarization on the resting MET current but independent from the presence or absence of Tmc2.

We then stimulated the hair bundle with 50 ms force steps to further investigate the dynamic adaptation properties of the MET current (Fig. 4). Upon deflecting the bundles in the excitatory direction, at −81 mV in the presence of 1.3 mM extracellular Ca2+, adaptation or time-dependent decline of the MET current occurred for small bundle displacements in OHCs from both Tmc2−/−:Tmc1+/+ and Tmc2−/−:Tmc1+Bth/Bth mice (Fig. 4A,B, left panels, respectively). Inhibitory hair bundle deflection shut off the small fraction of the current flowing at rest. All these manifestations of MET current adaptation were lost on stepping the membrane potential to +99 mV (Fig. 4A,B, right panels), a condition that prevents or strongly reduces Ca2+ entry via the MET channels. Stepping the membrane potential from −81 to +99 mV increased the resting P_open of the MET channel in OHCs from Tmc2−/−:Tmc1+/+ and Tmc2−/−:Tmc1+Bth/Bth mice (Fig. 4A,B, left), but not from Tmc2−/−:Tmc1+Bth/Bth mice (Fig. 4B,D), which is indicated by the lack of a leftward shift in the relationship between the normalized MET current and bundle displacement in the Beethoven mutant (Fig. 4D). The resting P_open at −81 mV was similar between Tmc1+/+, Tmc1Bth/Bth, and Tmc1+Bth/Bth (all were knock-outs for Tmc2; Fig. 4E, left). However, at +99 mV, the resting P_open in Tmc2−/−:Tmc1+/+ was significantly larger compared with both Tmc2−/−:Tmc1Bth/Bth and Tmc2−/−:Tmc1+Bth/Bth (p < 0.001, one-way ANOVA post hoc tests).

We further tested the effects of Ca2+ on the MET current of Beethoven mutants by comparing experiments in 1.3 mM Ca2+ and low extracellular Ca2+ (0.04 and 0.1 mM). These low Ca2+...
concentrations were selected because, although the bundles of adult mouse hair cells are surrounded by endolymph containing 0.02–0.04 mM Ca\(^{2+}\) (Bosher and Warren, 1978; Salt et al., 1989), immature cells are exposed to an estimated in vivo endolymphatic Ca\(^{2+}\) concentration in the low millimolar range (\(\leq 0.3\) mM; Johnson et al., 2012). In OHCs from both \(\text{Tmc1}^{+/+}\) and \(\text{Tmc1}^{\text{Bth/Bth}}\) mice, reducing the extracellular Ca\(^{2+}\) concentration from 1.3 to 0.1 mM (Fig. 5A, B, respectively) or 0.04 mM (data not shown) at \(-81\) mV increased the MET channel resting \(P_{\text{open}}\) as reported previously (Johnson et al., 2011; Corns et al., 2014). However, the overall dependence of \(P_{\text{open}}\) on extracellular Ca\(^{2+}\) was found to be significantly different between the two genotypes (\(p < 0.01\), two-way ANOVA; Fig. 5C–E), with the post hoc test analysis revealing a significant difference for only 0.1 mM Ca\(^{2+}\) between \(\text{Tmc1}^{+/+}\) and \(\text{Tmc1}^{\text{Bth/Bth}}\) (\(p < 0.001\)). Furthermore, although in \(\text{Tmc1}^{+/+}\) the \(P_{\text{open}}\) was similar between 0.1 and 0.04 mM Ca\(^{2+}\), in \(\text{Tmc1}^{\text{Bth/Bth}}\) it was significantly reduced in the former (\(p < 0.001\), one-way ANOVA). This points to an increased sensitivity of the open probability of the resting MET current to extracellular Ca\(^{2+}\) in \(\text{Tmc1}^{\text{Bth/Bth}}\). These findings differ from a recent report showing that the \(P_{\text{open}}\) of the MET current in Beethoven OHCs was only slightly affected when reducing the extracellular Ca\(^{2+}\) from 1.3 to 0.04 mM (Beurg et al., 2015), but within the range we have reported for the 0.1 mM Ca\(^{2+}\) solution (Fig. 5C–E). This discrepancy in Ca\(^{2+}\) sensitivity may have been attributable to incomplete exchange of the solution in the fluid jet used to stimulate the hair bundle when changing the different Ca\(^{2+}\) concentrations and genotypes.

Lowering the extracellular Ca\(^{2+}\) concentration also had the effect of increasing the size of the MET current (Fig. 5A, B), which stems from relief of the block by Ca\(^{2+}\) in the permeation pathway of the channel (Ricci and Fettiplace, 1998; Marcotti et al., 2005). For the experiments in which the MET current was recorded in 1.3 mM and either 0.1 or 0.04 mM extracellular Ca\(^{2+}\) from the same OHCs (see Materials and Methods), we found that the MET current size ratio (1.3/0.1 mM Ca\(^{2+}\) or 1.3/0.04 mM Ca\(^{2+}\)) recorded at \(-81\) mV was significantly smaller in \(\text{Tmc1}^{\text{Bth/Bth}}\) (1.3/0.1 mM Ca\(^{2+}\) = 0.5280 ± 0.0199, \(n = 9, p < 0.002; 1.3/0.04\) mM Ca\(^{2+}\) = 0.5871 ± 0.0218, \(n = 8, p < 0.001\)) than in \(\text{Tmc1}^{+/+}\) (1.3/0.1 mM Ca\(^{2+}\) = 1.38 ± 0.07, \(n = 23, p > 0.05; 1.3/0.04\) mM Ca\(^{2+}\) = 1.51 ± 0.09, \(n = 23, p > 0.05\)).
Ca\(^{2+}\) = 0.5153 ± 0.0178, \(n = 9, p < 0.02\) than in \(Tmc1^{+/+}\) OHCs (1.3/0.1 mM Ca\(^{2+}\) = 0.6976 ± 0.0422, \(n = 4\); 1.3/0.04 mM Ca\(^{2+}\) = 0.6106 ± 0.0285, \(n = 6\)). A smaller ratio in the mutants implies a stronger Ca\(^{2+}\) block of the MET current in the presence of 1.3 mM Ca\(^{2+}\) in \(Tmc1^{Bth/Bth}\) OHCs, which is in agreement with the reduced Ca\(^{2+}\) permeability and conductance observed in these cells, and all these findings indicate that the Beethoven point mutation directly affects the MET channel Ca\(^{2+}\) binding site located within the channel. A similar stronger Ca\(^{2+}\) block of the MET current has also been reported in Beethoven mouse IHCs (Pan et al., 2013).

Intracellular calcium modulation of MET currents in OHCs of Beethoven mice

We tested whether MET currents of \(Tmc1^{Bth/Bth}\) OHCs were directly regulated by free Ca\(^{2+}\) inside the stereocilia by changing the intracellular Ca\(^{2+}\) buffering capacity. We recorded MET currents at −81 mV in the presence of different concentrations of the fast Ca\(^{2+}\) buffer BAPTA (Fig. 6). In the presence of 0.1 mM BAPTA, nonsaturating bundle displacements caused the MET current to adapt in both genotypes (Fig. 6A, B, left panels), exactly as seen when 1 mM EGTA was used in the intracellular solution (Fig. 4). In the presence of 10 mM intracellular BAPTA, the time-dependent MET current decline was abolished and the resting \(P_{\text{open}}\), increased to near 50% of the maximal MET current in OHCs from both \(Tmc1^{+/+}\) (Fig. 6A, right) and \(Tmc1^{Bth/Bth}\) (Fig. 6B, right) mice. The relation between the MET current and bundle displacement (Fig. 6C, D) shows that increasing the intracellular BAPTA concentration from 0.1 to 10 mM significantly increased (\(p < 0.0001\)) the resting \(P_{\text{open}}\) of the MET current in both \(Tmc1^{+/+}\) (0.1 mM, 8.0 ± 1.6%, \(n = 4\); 10 mM, 39.6 ± 2.7%, \(n = 5\)) and \(Tmc1^{Bth/Bth}\) (0.1 mM, 10.4 ± 2.2%, \(n = 3\); 10 mM, 46.5 ± 9.9%, \(n = 6\)). No significant differences were seen between the two genotypes for both BAPTA concentrations. However, 3 and 5 mM BAPTA were less effective in shifting the MET current-bundle displacement curves in \(Tmc1^{Bth/Bth}\) than in \(Tmc1^{+/+}\) OHCs (Fig. 6C, D). In \(Tmc1^{+/+}\), increasing the BAPTA concentration from 0.1 mM to either 3 or 5 mM produced a highly significant increase in \(P_{\text{open}}\) (post hoc test from one-way ANOVA, \(p < 0.01\) and \(p < 0.001\), respectively; Fig. 6D); in \(Tmc1^{Bth/Bth}\), the same comparison produced no or a much reduced increase in \(P_{\text{open}}\) (n.s. and \(p < 0.05\), respectively; Fig. 6E).

Effects of DHS on the MET current in Beethoven mice

DHS is known to block the MET channel in hair cells of both nonmammals (Kroese et al., 1989; Ohmori, 1985; Ricci, 2002) and mammals, in which it has been shown to enter the cell by traversing the channel, i.e., it is a permeant blocker (Marcotti et al., 2005). The dependence of the DHS-induced channel block on extracellular Ca\(^{2+}\) and membrane potential (Kroese et al., 1989; Ricci, 2002; Marcotti et al., 2005) has led to the suggestion that the drug-binding site is within the permeation pathway of the
channel, and this binding site is accessible from both intracellular and extracellular sides of the plasma membrane (Marcotti et al., 2005; van Netten and Kros, 2007). MET currents were mainly recorded by stepping the membrane between -121 and +99 mV in 20 mV increments, although in some experiments, we extended the voltage range to between -161 and +159 mV. Using 50 Hz sinusoidal stimulation, we recorded the MET current from OHCs of both Tmc1+/+ and Tmc1Bth/Bth in the presence of dif-

Figure 7. The MET channel of OHCs of Beethoven mice has a reduced affinity to DHS. A, B, Block of saturating MET current by extracellular DHS in apical OHCs of Tmc1+/+ (A) and Tmc1Bth/Bth (B) mice in response to sinusoidal stimuli to the hair bundles (top) and at membrane potentials of -121 and +99 mV. Recordings were performed in the presence of 100 μM extracellular DHS. C, D, Average normalized MET current–voltage curves recorded from OHCs of P7–P8 Tmc1+/+ (C) and P6–P9 Tmc1Bth/Bth (D) for some of the DHS concentrations tested. Numbers of OHCs recorded under the different conditions were as follows: Tmc1+/+ control conditions (n = 16), 10 μM DHS (n = 15), 100 μM DHS (n = 14); Tmc1Bth/Bth control conditions (n = 7), 10 μM DHS (n = 7), 100 μM DHS (n = 6). E, Dose–response curves for the block of the MET current by extracellular DHS at -81 mV in OHCs from Tmc1+/+ (filled symbol) and Tmc1Bth/Bth (open symbol). Continuous lines are fits through the data using a Hill equation. Tmc1+/+ OHCs (P6–P8) half-blocking concentration of K

$\Delta E$ was 4.83 kT in Tmc1+/+ and 7.32 kT in Tmc1Bth/Bth, $\Delta E_b$ was -7.65 kT in Tmc1+/+ and -4.21 kT in Tmc1Bth/Bth.

$\Delta E_b$ was 4.83 kT in Tmc1+/+ and 7.32 kT in Tmc1Bth/Bth, $\Delta E_b$ was -7.65 kT in Tmc1+/+ and -4.21 kT in Tmc1Bth/Bth.

$\Delta E_b$ was 4.83 kT in Tmc1+/+ and 7.32 kT in Tmc1Bth/Bth, $\Delta E_b$ was -7.65 kT in Tmc1+/+ and -4.21 kT in Tmc1Bth/Bth.
different concentrations of extracellular (Fig. 7) and intracellular (Fig. 8) DHS. As described previously (Marcotti et al., 2005), extracellular DHS caused a voltage-dependent block of the MET current, pronounced at negative membrane potentials and relieved at positive potentials in both genotypes (Fig. 7A,B). The normalized current–voltage curves for the peak-to-peak MET current with and without extracellular DHS are shown in Figure 7, C (Tmc1+/+) and D (Tmc1Bth/Bth). The concentration for half block (Kp) of the MET current at −81 mV in Tmc1Bth/Bth (108 μM) was nearly eight times higher than that measured in Tmc1+/+ (14 μM; Fig. 7E). The voltage dependence of the block of the MET channel by DHS was investigated further by plotting the MET current in OHCs in the presence of the drug as a fraction of the control current (I/Is; Fig. 7F). The block of the MET current by DHS is partially relieved for values negative to approximately −80 mV in both genotypes. This behavior is consistent with DHS being pushed from its binding site and forced through the channel pore into the cytoplasm when sufficient electrical driving force is applied (Marcotti et al., 2005), indicating that DHS can still permeate the MET channel in Bth mutant mice. The fits through the data are according to the two-barrier one-binding-site model (Marcotti et al., 2005). Intracellular application of DHS inhibited the MET current at positive potentials in a similar dose-dependent manner in both Tmc1+/+ and Tmc1Bth/Bth OHCs (Fig. 8A–C) but, as described previously (Marcotti et al., 2005), with a reduced potency compared with extracellular DHS. In Tmc1Bth/Bth OHCs, the Kp for intracellular DHS block was 478 μM, which was nearly three times higher than that measured in for Tmc1+/+ OHCs (167 μM; Fig. 8D).

To calculate the absolute heights of the energy barriers and to obtain an indication of whether the Beethoven mutation affects the rate of entry of DHS molecules into the OHCs, we measured the time constant of the initial decline of the MET current in the presence of the antibiotic (Marcotti et al., 2005). For these experiments, hair bundles were initially deflected toward the inhibitory barriers, but the intracellular barrier site in the channel pore, and raises the heights of both energy barriers, but the intracellular barrier Ei. The number of DHS molecules entering into the OHCs was estimated as before by assuming a driving force of −150 mV, a total of 80 MET channels per OHC and a resting open probability of the MET channel of 0.3 (Marcotti et al., 2005), but using 1.3 ms extracellular Ca2+ as that was how the present experiments were conducted. The result was a substantially reduced entry of DHS molecules in the Beethoven mutants (Tmc1+/+, 2232/s; Tmc1Bth/Bth, 335/s).

Figure 8. Reduced block by intracellular DHS in OHCs of Beethoven mice. A, Block of MET current by intracellular DHS in apical P7 OHCs of Tmc1+/+ (left) and Tmc1Bth/Bth (right) mice at membrane potentials of −121 and +99 mV in the presence of 500 μM DHS. Experimental protocol as described in Figure 7. B, C, Average normalized MET current–voltage curves from OHCs of P7–P8 Tmc1+/+ (B) and P6–P9 Tmc1Bth/Bth (C) for some of DHS concentrations tested. Note that intracellular DHS blocks the MET current at depolarized potentials in OHCs from both genotypes. Numbers of OHCs recorded under the different conditions were as follows: Tmc1+/+ control conditions (n = 16), 100 μM DHS (n = 5), 500 μM DHS (n = 8), and 10 mM DHS (n = 6); Tmc1Bth/Bth control conditions (n = 7), 100 μM DHS (n = 6), 500 μM DHS (n = 13), and 10 mM DHS (n = 6). D, Dose–response curves for the block of the MET current by intracellular DHS at +119 mV in OHCs from Tmc1+/+ (filled symbols) and Tmc1Bth/Bth (open symbols). Continuous lines are fits through the data using a Hill equation. Tmc1+/+ OHCs (P7–P8) half-blocking concentration of Kp of 167 ± 40 μM and a Hill coefficient nH of 1.1 ± 0.3 (number of OHCs from left to right: 5, 5, 8, 6; Tmc1Bth/Bth OHCs (P6–P9) half-blocking concentration of Kp of 479 ± 104 μM and a Hill coefficient nH of 0.9 ± 0.2 (number of OHCs from left to right: 3, 4, 5, 13, 5, 4, 6).
Discussion
In this study, we determined that the M412K point mutation in TMC1 of *Beethoven* mice leads to a reduced Ca\(^{2+}\) permeability, more so in *Tmc1\(^{Bth/Bth}\)* than in *Tmc1\(^{Bth/+}\)*, and conductance of the MET channel of OHCs. We also found that the M412K mutation causes a reduced affinity for the MET channel blocker DHS, which was more pronounced when the aminoglycoside was applied extracellularly. The effects on both Ca\(^{2+}\) permeability and DHS block are likely direct consequences of the replacement of a neutral methionine residue by a positively charged lysine affecting the permeation pathway of the channel.

The *Bth* mutation also affects the sensitivity of the MET channel to Ca\(^{2+}\) as manifested by the altered shifts in the current-displacement curves on manipulation of extracellular and intracellular Ca\(^{2+}\). In lower vertebrates, these shifts have been interpreted as steady-state manifestations of Ca\(^{2+}\)-dependent adaptation (Assad et al., 1989; Crawford et al., 1991; Ricci et al., 1998). Recently, it has been proposed that this is not the case for mammalian cochlear hair cells (Peng et al., 2013), but this view has been challenged on the grounds of technical issues because of glass-probe bundle stimulation (Corns et al., 2014) and underestimation of the Ca\(^{2+}\) concentrations that can be reached at the intracellular face of the MET channel (Beurg et al., 2015). In our view, the different effects caused by the *Bth* mutation can be most coherently explained by assuming that it alters Ca\(^{2+}\)-dependent adaptation, so that adaptation is not wild-type controls. Taking 0.04 mM extracellular Ca\(^{2+}\) as the reference condition, with little adaptation (Corns et al., 2014), increasing extracellular Ca\(^{2+}\) to 0.1 mM induced a stronger rightward adaptive shift in the current-displacement relation of OHCs of *Bth* mutant mice than in controls. Conversely, taking 0.1 mM intracellular BAPTA as the reference for intracellular Ca\(^{2+}\) buffering, 3 mM BAPTA was less able to induce a leftward shift in the OHCs of *Bth* mutants. It appears that, in *Bth* mutants, the sensitivity to Ca\(^{2+}\) at the sensor for adaptation, which is either part of the intracellular side of the MET channel or positioned very near it (Wu et al., 1999), is, surprisingly, increased, most likely as a downstream compensation mechanism for reduced Ca\(^{2+}\) entry through the MET channel. Our findings show that the *Bth* mutation affects Ca\(^{2+}\) permeability, the Ca\(^{2+}\) and DHS binding site within the channel and Ca\(^{2+}\)-dependent adaptation, indicating that TMC1 is a crucial component of the MET channel in hair cells.

The position of the *Beethoven* mutation
The topology of TMC1 is currently unknown, mainly because the protein is retained in the endoplasmic reticulum (ER) when using in vitro conditions (Kawashima et al., 2011). Using antibodies directed against epitope-tagged versions of the human TMC1 expressed in the ER of heterologous cell lines, TMC1 was predicted to have six transmembrane domains (Labay et al., 2010). If the same transmembrane topology of TMC1 is maintained in the plasma membrane, then the *Bth* M412K point mutation, which is predicted to be in the extracellular loop between the third and fourth membrane spanning domains (Holt et al., 2014), is likely to be located extracellularly. However, the number of transmembrane domains for TMC1 in vivo is still unknown, with some algorithms predicting the presence of up to four additional transmembrane regions (Kurima et al., 2002; Keresztes et al., 2003). Therefore, the exact location of M412K will depend on how many of the additional transmembrane domains are present in the native TMC1. Evidence for Tmci forming a channel comes from the heterologous expression of a *Caenorhabditis elegans* homolog of TMC1 (Chatzigeorgiou et al., 2013), which is critical for salt sensation and was inhibited by the stretch-activated channel blocker gadolinium, reported to block the hair cell MET channel (Kimitsu et al., 1996).

Block of the OHC MET channel by DHS can be described by a two-barrier one-binding-site model with the binding site positioned at a relative electrical distance of 0.79 into the channel from the extracellular side. The binding site is located between 0.045 and 0.79, as measured across the membrane from the extracellular side. The two barriers sandwich the binding site for DHS at a relative electrical distance of 0.79 into the channel from the extracellular side. The two barriers sandwich the binding site for DHS at a relative electrical distance of 0.79 with a minimum in free energy, \(E_f\) of \(-7.65\) kT for controls and \(-4.21\) kT for *Bth*.

**Figure 9.** DHS entry into *Beethoven* OHCs. **A, B,** Block of saturating MET current by extracellular DHS in apical P6 OHCs of *Tmc1\(^{+/+}\)* (A) and *Tmc1\(^{Bth/Bth}\)* (B) mice in response to step stimuli to the hair bundles (top) and at membrane potentials of \(-81\) mV. MET channels were first closed by inhibitory bundle displacement and then fully opened by an excitatory deflection. Recordings were performed before and during the application of the antibiotic (see Results). The decay of the MET current in the presence of DHS was fitted using a single exponential (gray). **C, D,** Rate-constant \(k_r\) was obtained by investigating the DHS-binding kinetics as described previously (Marcotti et al., 2005). The inverse of the time constant of binding (1/\(k_r\)), which was obtained by the experiments shown in A and B, was plotted against two different extracellular DHS concentrations in each genotype. Solid lines indicate the fits and the slope constant of binding (1/

\[
\text{Energy profiles of two-barrier one-binding-site model for the MET-channel pore of TMC1}\].
two energy barriers, with the intracellular barrier being higher than that on the extracellular side, which accounts in part for the much higher blocking potency of DHS when applied extracellularly (Marcotti et al., 2005; van Netten and Kros, 2007). The presence of a wide channel vestibule facing the extracellular side allows a rapid almost diffusion-limited entry of blockers into its pore region (van Netten and Kros, 2007). The *Beethoven* mutation caused the MET channel to be less sensitive to block by DHS, with a greater reduction occurring when the aminoglycoside was applied extracellularly (7.5-fold) than intracellularly (3.0-fold). The mutation lowered the affinity of the DHS binding site by 3.34 $kT$. The height of the intracellular energy barrier $E_2$ was increased by 4.39 $kT$, hindering movement in either direction of DHS over this barrier, whereas the extracellular barrier $E_1$ was increased to a lesser extent, by 1.90 $kT$ (Fig. 9). This agrees with the positively charged lysine of the *Bth* mutation most strongly affecting the intracellular side of the channel pore (Fig. 10), near the binding site, at 0.79 electrical distance, and the intracellular barrier. This lysine is thus likely to be present in or near the permeation pore of the MET channel, in which it reduces the affinity of DHS for a binding site for polycations deep in the vestibule of the channel, near the selectivity filter of the narrow pore region (Van Netten and Kros, 2007).

It is interesting to compare the reduced affinity for DHS of OHCs of *Bth* mutant mice with the anomalous MET currents recorded from cochlear hair cells of mice that have no tip-link-gated MET channels (Marcotti et al., 2014), in which the $K_D$ was, at 894 $\mu$m, eight times higher than in the *Tmc1Bth/Bth* OHCs. There, it was proposed that the vestibule, and hence the polycationic binding site, was lacking. Together, our findings strengthen the evidence for TMC1 being an integral component of the MET channel (Holt et al., 2014). In the mechanosensitive channel Piezo1, an E2133K point mutation (replacing a negatively charged glutamate with a positive lysine) similarly reduced its Ca$^{2+}$ permeability and the blocking potency of the polycationic ruthenium red, leading to the conclusion that this residue must be at or near the pore (Coste et al., 2015).

**Functional consequences of the Beethoven mutation**

The size of the MET currents in *Beethoven* mutant OHCs is normal at least until around the onset of hearing, and yet compound action potential (CAP) responses are absent in young mice (Marcotti et al., 2006). This suggests that a more subtle defect is responsible for the progressive hearing loss phenotype in the *Beethoven* mutant mice (Vreugde et al., 2002; Marcotti et al., 2006). Recently, it has been proposed that the reduced expression of the plasma membrane Ca$^{2+}$ ATPase 2 (PMCA2) pump in the stereociliary bundle of *Beethoven* OHCs could contribute to the hearing loss (Beurg et al., 2015). Defects in PMCA2 have been linked to deafness in mice (Street et al., 1998; Bortolozzi et al., 2010) and associated with reductions in endocochlear potential (Wood et al., 2004) and endolympathic Ca$^{2+}$, which would disrupt the PCDH15–cadherin 23 interaction at the tip link (Kazmierczak et al., 2007). However, given that both endocochlear potential (Marcotti et al., 2006) and MET current size (Fig. 1) in *Beethoven* mice were normal, reduced PMCA2 is unlikely to contribute significantly to the hearing phenotype in *Bth* mutants.

We propose that the stronger MET current adaptation is the main causative factor of the hearing loss. The hair bundles of the immature mouse cochlea OHCs are exposed to an *in vivo* endolympathic Ca$^{2+}$ concentration of $\leq$0.3 mM but higher than the 0.04 mM in the mature cochlea (Johnson et al., 2012). Despite the reduced Ca$^{2+}$ influx into the MET channel of *Bth* mutants, the channel is kept in a more strongly adapted state, already evident at 0.1 mM Ca$^{2+}$, which results in a lower fraction of the current activated at the resting bundle position. The reduced resting open probability of the immature hair bundle of *Bth* mutant OHCs will reduce the standing inward MET current and cause their membrane potential ($V_m$) to hyperpolarize compared with control cells (approximately $-40$ mV; Johnson et al., 2011). The similarity between our findings in OHCs and those reported previously in IHCs (Pan et al., 2013) suggests that the *Bth* mutation is also likely to hyperpolarize the IHC resting $V_m$ (approximately $-60$ mV; Johnson et al., 2011). Alteration of the normal electrical activity in developing cochlear hair cells has been associated with defects in the synaptic machinery and basolateral membrane properties (Roux et al., 2009; Johnson et al., 2013), which resemble those observed in the *Bth* mutant hair cells (Marcotti et al., 2006). The reduced expression of the potassium current $I_{K,Na}$ from approximately $P8$ onward in OHCs (Vreugde et al., 2002; Marcotti et al., 2006) is likely to contribute to the hearing defects observed in young and adult mice (Marcotti et al., 2006). Although *Bth* mutant OHCs retain their characteristic electromotile activity (Marcotti et al., 2006), the shift in their $V_m$ will affect the optimal activation of the motor protein prestin (Ashmore, 2008; Johnson et al., 2011), which drives electromotility (He et al., 1994; Marcotti and Kros, 1999).
References
Ashmore J (2008) Cochlear outer hair cell motility. Physiol Rev 88:173–210. CrossRef Medline
Assad JA, Hacohen N, Corey DP (1989) Voltage dependence of adaptation and active bundle movement in bullfrog saccular hair cells. Proc Natl Acad Sci U S A 86:2918–2922. CrossRef Medline
Assad JA, Shepherd GM, Corey DP (1991) Tip-link integrity and mechanical transduction in vertebrate hair cells. Neuron 7:985–994. CrossRef Medline
Beurg M, Fettipalce R, Nam JH, Ricci AJ (2009) Localization of inner hair cell mechanotransducer channels using high-speed calcium imaging. Nat Neurosci 12:553–558. CrossRef Medline
Beurg M, Kim KK, Fettipalce R (2014) Conductance and block of hair-cell mechanotransducer channels in transmembrane channel-like protein mutants. J Gen Physiol 144:55–69. CrossRef Medline
Beurg M, Goldberg AR, Fettipalce R (2015) The effects of Tmc1 Beethoven mutation on mechanotransducer channel function in cochlear hair cells. J Gen Physiol 146:233–243. CrossRef Medline
Bortolozzi M, Brini M, Parkinson N, Crispino G, Scimemi P, Di Siati RD, Di Leva F, Parker A, Ortolano S, Azlian E, Brown SD, Carafoli E, Mannamo F (2010) The novel PMC2A pump mutation Tommy impairs cytosolic calcium clearance in hair cells and links to deafness in mice. J Biol Chem 285:37693–37703. CrossRef Medline
Bosher SK, Warren RL (1978) Very low calcium content of cochlear ensolymph, an extracellular fluid. Nature 273:377–378. CrossRef Medline
Chatzigorgeou M, Bang S, Hwang SW, Schafer WR (2013) tmc-1 encodes a sodium-sensitive channel required for salt chemosensation in C. elegans. Nature 494:95–99. CrossRef Medline
Corns LF, Johnson SL, Kros CJ, Marcotti W (2014) Calcium entry into steroeola drives adaptation of the mechanoelectrical transducer current of mammalian cochlear hair cells. Proc Natl Acad Sci U S A 111:14918–14923. CrossRef Medline
Coste B, Murthy SE, Mathur J, Schmidt M, Mechioukhi Y, Delmas P, Patapoutian A (2015) Piezo1 ion channel pore properties are dictated by G-terminal region. Nat Commun 6:7223. CrossRef Medline
Crawford AC, Evans MG, Fettipalce R (1991) The actions of calcium on the mecano-electrical transducer current of turtle hair cells. J Physiol 434:369–398. CrossRef Medline
Goodyear RJ, Marcotti W, Kros CJ, Richardson GP (2005) Development and properties of stereociliary link type in hair cells of the mouse cochlea. J Comp Neurol 485:75–85. CrossRef Medline
He DZ, Evans BN, Dallos P (1994) First appearance and development of mammalian cochlear inner hair cells. J Neurosci 14:3392–3405. CrossRef Medline
Holt JR, Pan B, Koussa MA, Asai Y (2014) TMC function in hair cell transduction. Hear Res 311:17–24. CrossRef Medline
Johnson SL, Beurg M, Marcotti W, Fettipalce R (2011) Prestin-driven cochlear amplification is not limited by the outer hair cell membrane time constant. Neuron 70:1143–1154. CrossRef Medline
Johnson SL, Kennedy HJ, Holley MC, Fettipalce R, Marcotti W (2012) The resting transducer current drives spontaneous activity in pre-hearing mammalian cochlear inner hair cells. J Neurosci 32:10479–10483. CrossRef Medline
Johnson SL, Kuhn S, Franz C, Ingham N, Furness DN, Knipper M, Steel KP, Adelman JP, Holley MC, Marcotti W (2013) Presynaptic maturation in auditory hair cells requires a critical period of sensory-independent spiking activity. Proc Natl Acad Sci U S A 110:8720–8725. CrossRef Medline
Kawashima Y, Gélcós GS, Kurima K, Labay V, Lelli A, Asai Y, Makishima T, Wu DK, Della Santina CC, Holt JR, Griffith AJ (2011) Mechanotransduction in mouse inner ear hair cells requires transmembrane channel-like genes. J Clin Invest 121:4796–4809. CrossRef Medline
Kawashima Y, Gélcós GS, Kurima K, Labay V, Lelli A, Asai Y, Makishima T, Wu DK, Della Santina CC, Holt JR, Griffith AJ (2011) Mechanotransduction in mouse inner ear hair cells requires transmembrane channel-like genes. J Clin Invest 121:4796–4809. CrossRef Medline
Kazmierczak P, Sakaguchi H, Tokita J, Wilson-Kubalek EM, Milligan RA, Müller U, Kachar B (2007) Cadherin 23 and protocadherin 15 interact to form tip-link filaments in sensory hair cells. Nature 449:87–91. CrossRef Medline
Keresztes G, Mutai H, Heller S (2003) TMC and EVER genes belong to a larger novel family, the TMC gene family encoding transmembrane proteins. BMC Genomics 4:24. CrossRef Medline
Kimitsu T, Nakagawa T, Hisaki K, Komune S, Komiyama S (1996) Gadolinium blocks mechano-electric transducer current in chick cochlear hair cells. Hear Res 101:75–80. CrossRef Medline
Kroese AB, Das A, Hudspeth AJ (1989) Blockage of the transduction channels of hair cells in the bullfrog’s saccule by aminoglycoside antibiotics. Hear Res 37:203–217. CrossRef Medline
Kros CJ, Rüsch A, Richardson GP (1992) Mechano-electrical transducer currents in hair cells of the cultured neonatal mouse cochlea. Proc Biol Sci 249:185–193. CrossRef Medline
Kurima K, Peters LM, Yang Y, Riazuddin S, Naz S, Arnaud D, Drury S, Mo J, Makishima T, Ghosh M, Menon PS, Deshmukh D, Oddoux C, Oster H, Khan S, Riazuddin S, Deininger PL, Hampton LL, Sullivan SL, Battey JF Jr, Keats BJ, Wilcox ER, Friedman TB, Griffith AJ (2002) Dominant and recessive deafness caused by mutations of a novel gene, TMC1, required for cochlear hair-cell function. Nat Genet 30:277–284. CrossRef Medline
Kurima K, Ebrahim S, Pan B, Sedlavec M, Sengupta P, Mills BA, Cui R, Nakaniishi H, Fujikawa T, Kawashima Y, Choi BY, Monahan K, Holt JR, Griffith AJ, Kachar B (2015) TMC1 and TMC2 localize at the site of mechatransduction in mammalian inner ear hair cell stereocilia. Cell Rep 12:1606–1617. CrossRef Medline
Labay V, Weichert RM, Makishima T, Griffith AJ (2010) Topology of transmembrane-channel-like gene 1 protein. Biochemistry 49:8592–8598. CrossRef Medline
Marcotti W, Kros CJ (1999) Developmental expression of the potassium current IK, Ni contributes to maturation of the mouse outer hair cells. J Physiol 520:653–660. CrossRef Medline
Marcotti W, van Netten SM, Kros CJ (2005) The aminoglycoside antibiotic dihydrostreptomycin rapidly enters hair cells through the mecano-electrical transducer channels. J Physiol 567:505–521. CrossRef Medline
Marcotti W, Erven A, Johnson SL, Steel KP, Kros CJ (2006) Tmc1 is necessary for normal functional maturation and survival of inner and outer hair cells in the mouse cochlea. J Physiol 574:677–698. CrossRef Medline
Marcotti W, Corns LF, Desmonds T, Kirkwood NK, Richardson GP, Kros CJ (2014) Transduction without tip links in cochlear hair cells is mediated by ion channels with permeation properties distinct from those of the mecano-electrical transducer channel. J Neurosci 34:500–514. CrossRef Medline
Nakanishi H, Kurima K, Kawashima Y, Griffith AJ (2014) Mutations of TMC1 cause deafness by disrupting mechanoelectrical transduction. Audnasus Larynx 41:399–408. CrossRef Medline
Nam JH, Peng AW, Ricci AJ (2015) Underestimated sensitivity of mammalian cochlear hair cells due to splay between stereociliary columns. Biophys J 108:2633–2647. CrossRef Medline
Ohmori H (1985) Mechano-electrical transduction currents in isolated vestibular hair cells of the chick. J Physiol 359:189–217. CrossRef Medline
Pan B, Gélcós GS, Asai Y, Horwitz GC, Kurima K, Ishikawa K, Kawashima Y, Griffith AJ, Holt JR (2013) TMC1 and TMC2 are components of the mechatransduction channel in hair cells of the mammalian inner ear. Neuron 79:504–515. CrossRef Medline
Partanen JI (2010) Re-evaluation of the thermodynamic activity quantities in aqueous rubidium and cesium chloride solutions at 25°C. J Chem Eng Data 55:249–257. CrossRef Medline
Peng AW, Effertz T, Ricci AJ (2013) Adaptation of mammalian auditory hair cell mechatransduction is independent of calcium entry. Neuron 80:960–972. CrossRef Medline
Pickles JO, Comis SD, Osborne MP (1984) Cross-links between stereocilia in the guinea pig organ of Corti, and their possible relation to sensory transduction. Hear Res 15:103–112. CrossRef Medline
Rard JA, Clegg SL (1997) Critical evaluation of the thermodynamic properties of aqueous calcium chloride. I. Osmotic and activity coefficients of 0–10.77 mol.L−1 aqueous calcium chloride solutions at 298.15 K and correlation with extended pitzer ion-interaction models. J Chem Eng Data 42:819–849. CrossRef Medline
Ricci A (2002) Differences in mecano-transducer channel kinetics underlie tonotopic distribution of fast adaptation in auditory hair cells. J Neurophysiol 87:1738–1748. Medline
Ricci AJ, Fettipalce R (1998) Calcium permeation of the turtle hair cell mechano-transducer channel and its relation to the composition of endolymph. J Physiol 506:159–173. CrossRef Medline
