Transduction without Tip Links in Cochlear Hair Cells Is Mediated by Ion Channels with Permeation Properties Distinct from Those of the Mechano-Electrical Transducer Channel

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Introduction

The tip links of sensory hair cells are composed of two members of the cadherin superfamily of cell–cell adhesion molecules, protocadherin 15 (PCDH15) and cadherin 23 (CDH23) (Sollner et al., 2004; Siemens et al., 2004; Ahmed et al., 2006). PCDH15 is located at the lower end of the tip link, whereas CDH23 is located at the upper end (Kazmierczak et al., 2007). The mechanoelectrical-transducer (MET) channels are localized at the tips of the shorter stereocilia at the lower end of each link (Beurg et al., 2009). One would therefore expect hair cells in which PCDH15 or CDH23 are absent to lack MET currents. Unusual MET currents with abnormal directional sensitivity have, however, been observed in outer hair cells (OHCs) of Pcdh15 and Cdh23 mutant mice, suggesting MET channels can be gated by means other than via tip links (Alagramam et al., 2011). By examining the characteristics of MET currents that can be elicited under a variety of conditions in cochlear hair cells that lack normal tip links, we show that this phenomenon is not restricted to hair cells with mutations in tip-link proteins. Our results reveal that the pore properties of the underlying ion channels are different from those of the normal MET channels of cochlear hair cells.

Materials and Methods

OHCs and inner hair cells (IHCs) from CD-1 and C57BL/6J mice of either sex were studied from embryonic day (E)18.5 to postnatal day (P)10 where the day of birth (P0) corresponds to E19.5. Recordings were performed from acutely dissected organs of Corti (Marcotti et al., 2003) or in organotypic cultures (Russell and Richardson, 1987) that had been maintained for 1–3 d in vitro. Most recordings were made from hair cells situated in the apical coil. When recordings from basal-coil hair cells are reported this is stated explicitly. Some recordings were performed from outer hair cells (OHCs) from homozygous Myo7a<sup>−/−</sup>, Myo7a<sup>−/−</sup>Cdhi15<sup>−/−</sup>, Cdhi23<sup>−/−</sup>, and Pcdh15<sup>−/−</sup> mutant mice (Kros et al., 2002; Alagramam et al., 2011). For experiments with embryos, mice were paired overnight and checked for vaginal plugs the following morning. Assuming ovulation occurs midway through the

Tip links between adjacent stereocilia are believed to gate mechanoelectrical transducer (MET) channels and mediate the electrical responses of sensory hair cells. We found that mouse auditory hair cells that lack tip links due to genetic mutations or exposure to the Ca<sup>2+</sup> chelator BAPTA can, however, still respond to mechanical stimuli. These MET currents have unusual properties and are predominantly of the opposite polarity relative to those measured when tip links are present. There are other striking differences, for example, the channels are usually all closed when the hair cell is not stimulated and the currents in response to strong stimuli can be substantially larger than normal. These anomalous MET currents can also be elicited early in development, before the onset of mechanoelectrical transduction with normal response polarity. Current–voltage curves of the anomalous MET currents are linear and do not show the rectification characteristic of normal MET currents. The permeant MET channel blocker dihydrostreptomycin is two orders of magnitude less effective in blocking the anomalous MET currents. The findings suggest the presence of a large population of MET channels with pore properties that are distinct from those of normal MET channels. These channels are not gated by hair-bundle links and can be activated under a variety of conditions in which normal tip-link-mediated transduction is not operational.

Key words: aminoglycoside antibiotics; cochlea; hair cell; mechanoelectrical transduction

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dark cycle, the mid-point of the light cycle of the day following mating is considered to be E0.5. Adult and neonatal mice were killed by cervical dislocation and embryos by decapitation, in accordance with UK Home Office regulations.

For the experiments using acutely isolated organs of Corti, cochleae were dissected in normal extracellular solution (in mM): 135 NaCl, 5.8 KCl, 1.3 CaCl2, 0.9 MgCl2, 0.7 NaH2PO4, 5.6 D-glucose, 10 HEPES-NaOH. Sodium pyruvate (2 mM), MEM amino acids solution 5.6 D-glucose, 10 HEPES-NaOH. Sodium pyruvate (2 mM), MEM amino acids solution (50 ×, without L-glutamine), and MEM vitamins solution (100 ×) were added from concentrates (Fisher Scientific). The pH was adjusted to 7.5 (osmolality ~308 mmol kg⁻¹).

The dissected organs of Corti were transferred to a microscope chamber and immobilized using a nylon mesh fixed to a stainless steel ring. Where organotypic cultures were used, they were transferred to the microscope chamber on their collagen substrate and did not require further immobilization. The chamber was continuously perfused with the above extracellular solution. The organs of Corti were observed with an upright microscope (Zeiss, Leica) with Nomarski differential interference contrast optics (40 × or 63 × water-immersion objectives).

Whole-cell patch-clamp recordings were performed at room temperature using EPC-8 (HEKA) or Optopatch (Cairn Research) patch-clamp amplifiers. Patch pipettes (2–3 MΩ) contained the following (in mM): 135 CsCl (or 106 Cs-glutamate, 20 CsCl), 2.5 MgCl2, 1 EGTA-CsOH, 2.5 Na2ATP, 10 sodium phosphocreatine, 5 CsCl (or 106 Cs-glutamate, 20 CsCl), 2.5

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**Figure 1.** MET currents of OHCs before and during BAPTA treatment. A, B, Currents in response to sinusoidal force stimuli recorded from a P3 OHC of a wild-type CD-1 mouse before (A) and during (B) superfusion with 5 mM BAPTA. Membrane potential was stepped between −84 and +76 mV in 20 mV increments from a holding potential of −84 mV. For clarity only every other voltage level is shown. The driver voltage (DV: 45 Hz sinusoid, 25 V amplitude) to the fluid jet is shown above the traces. C, MET current recorded from the same OHC shown in A and B 2 min after the start of BAPTA application and after moving the fluid jet very close to the bundle until anomalous currents were seen. Large currents (note different scale of ordinate) were elicited during negative stimuli. D, Peak-to-peak–voltage curves obtained from the MET currentsshown in A–C. Fits to single-energy-barrier model (see Materials and Methods) are as follows: control k = 73 pA, Vr = +5.5 mV, Vh = 27 mV, and γ = 0.45; 2 min after BAPTA k = 1047 pA, Vr = +3.1 mV, Vh = 50 mV, and γ = 0.47.

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For the experiments investigating the pore properties of the channels, the aminoglycoside antibiotic dihydrostreptomycin (DHS) was locally superfused at concentrations ranging from 10 μM to 100 μM. Dose–response curves were fitted with the Hill equation:

\[
\frac{I_{\text{DHS}}}{I_c} = \frac{1}{1 + \left(\frac{[D]}{K_d}\right)^n}
\]

where \(I_{\text{DHS}}\) is the current in the presence of DHS, \(I_c\) is the control current, \(K_d\) is the dissociation constant, and \([D]\) is the concentration of DHS and \(n\) is the Hill coefficient. Stocks of DHS (molecular weight 730.7) were prepared at 10 μM (for final concentrations up to and including 300 μM) and 100 μM. Tip links were severed by bath perfusion of extracellular solution containing 5 μM BAPTA for 5 min. In a few experiments, neomycin (molecular weight 908.9) was used to study MET current block of homozygous Mys076 mutants.

The fits through the current–voltage curves are according to a single-energy-barrier model:

\[
I(V) = k \left( \exp \left( \frac{1 - \gamma(V - V_r)}{V_s} \right) - \exp \left( -\frac{\gamma(V - V_r)}{V_s} \right) \right)
\]

where \(k\) is a proportionality constant, \(V_r\) is the reversal potential, \(V_s\) is a measure for the steepness of the rectification, and \(\gamma\) is the fractional distance within the membrane’s electrical field of an energy barrier, as measured from the outside (Kros et al., 1992), or to a straight line, as appropriate. The size of the developing MET currents was fitted with a sigmoidal growth curve.

\[
I = \frac{I_{\text{max}}}{1 + \exp \left( -\frac{t - t_{1/2}}{S} \right)}
\]
Figure 2. MET currents of IHCs before and during BAPTA treatment. A, Saturating MET currents recorded from a P6 IHC of a wild-type C57BL/6J mouse, at a membrane potential of −81 mV. The driver voltage (DV: 50 Hz sinusoid, 40 V amplitude) to the fluid jet is shown above the current trace. B, No MET currents were recorded upon superfusion of 5 mM BAPTA. C, MET currents recorded from the same IHC shown in B 4 min after the start of BAPTA application and after moving the fluid jet very close to the bundle until anomalous currents were seen. Anomalous currents were elicited during negative stimuli. Arrows point to peak of negative phase of first sinusoid.

Figure 3. MET currents in response to orthogonal hair-bundle stimulation. A, B, Saturating MET currents recorded from a P7 OHC (A) and a P5 IHC (B) of wild-type C57BL/6J mice, at a membrane potential of −81 mV. The fluid jet (DV: 50 Hz sinusoid, 40 V amplitude, top) was positioned orthogonally to the axis of mechanical sensitivity of the hair bundle. C, D, MET currents recorded from a P7 OHC (C) and a P6 IHC (D) −5 min after the start of BAPTA application and after moving the fluid jet very close to the bundle until anomalous currents were seen.

where \( t_{1/2} \) is the age at which half the maximal current is reached and \( s \) is the slope, a measure for the steepness of growth. Means are reported as ±SEM in text and figures.

Results

MET currents in OHCs and IHCs after cleaving tip links with BAPTA

Exposure of hair bundles to the Ca\(^{2+}\) chelator BAPTA cuts both the tip links and the kinocilial links (Assad et al., 1991; Crawford et al., 1991; Gooyear and Richardson, 2003). Before the application of 5 mM BAPTA, a sinusoidal force stimulus from a fluid jet in the positive direction (toward the kinocilium) elicited large MET currents in wild-type OHCs from CD-1 or C57BL/6J mice, whereas negative stimuli closed the fraction of the MET current activated at rest (Fig. 1A). The currents reversed near 0 mV and the resting MET current increased at positive potentials, presumably due to a reduction in Ca\(^{2+}\)-dependent adaptation (Crawford et al., 1989). Upon application of 5 mM BAPTA the MET currents briefly increased in size in some cells (from −670 to −1380 pA in one OHC, and from −883 to −1630 pA in another, both at −124 mV; data not shown) due to a reduction of the permeant block by Ca\(^{2+}\) ions in the channel pore, before disappearing within 10 s (Fig. 1B). On applying strong mechanical stimulation in the continuing presence of BAPTA, mechanosensitive currents were elicited that could be larger than normal MET currents (Fig. 1C,D), up to −2 nA at −84 mV. The currents were not activated at the resting position of the bundle and occurred preferentially in response to negative force stimuli. During a recording from any given cell, currents were quite variable in size, likely due to the difficulty of applying strong enough mechanical stimulation, and in their directional sensitivity (see below).

In P6–P9 IHCs (\( n = 6 \)) BAPTA treatment of normally transducing cells (Fig. 2A) also resulted in anomalous MET currents (recorded in the continuing presence of BAPTA) that were elicited during the negative phase of the sine-wave stimulus (−464 ± 146 pA at −84 mV, range −65 to −1140 pA, \( n = 6 \)) following a period in which no MET current could be evoked (Fig. 2B,C). The size of the anomalous MET current was comparable to, but more variable than, the size of the normal MET current in control P6–P9 IHCs (−642 ± 33 pA at −84 mV, range −525 to −708 pA, \( n = 5 \)). In contrast to the anomalous current seen in OHCs, two of six IHCs recorded from in the presence of BAPTA (Fig. 2C) had a small resting current (1.6 ± 1.1% of the maximum current) at −84 mV. All five control IHCs had a resting MET current (3.3 ± 1.6%).

It was of interest to establish whether the anomalous MET currents were related at all to the axis of mechanical sensitivity of the hair bundle. We investigated this by stimulating the hair bundle of some OHCs and IHCs with the fluid jet positioned orthogonally to this axis before and during superfusion with 5 mM BAPTA. The results were not entirely straightforward because, even before BAPTA, inward MET currents were observed in OHCs (Fig. 3A) and IHCs (Fig. 3B) both for fluid flowing out of (positive-driver voltage) and into (negative-driver voltage) the jet. These bidirectional responses were presumably due to both halves of the V-shaped (OHCs) or more rounded (IHCs) bundle responding to the vectorial force directed along the axis toward
the kinocilium. During application of BAPTA the anomalous MET current could be observed, to a variable extent, in both phases of the stimulus for OHCs and IHCs (Fig. 3C,D, respectively). The maximum size of the anomalous MET currents recorded at −84 mV with orthogonal bundle stimulation during BAPTA application was −416 ± 139 pA for three OHCs and −430 ± 168 pA for four IHCs. The finding of bidirectional anomalous MET currents in response to orthogonal stimulation, while not conclusive, is at least consistent with the gating of the anomalous MET currents being indeed related to the axis of mechanical sensitivity of normal, tip-link gated MET currents, but with responses predominantly in the opposite direction.

**Gating and kinetics of anomalous MET currents**

These anomalous MET currents recorded during or after BAPTA application closely resembled MET currents reported in homozygous Cdhl23−/− and Peldh15−/− mutant mice that lack tip links (Alagramam et al., 2011) and in Myo7a−/− mutant mice lacking functional myosin VIIA (MYO7A) (Kros et al., 2002). Indeed, we still observed large anomalous MET currents in OHCs of homozygous Cdhl23−/− mutants in the continuous presence of 5 mM extracellular BAPTA (Fig. 4A,B). Similar findings were made in an additional three OHCs. We also found that the currents of OHCs from homozygous Myo7a−/− mice (n = 6) were resistant to treatment with 5 mM BAPTA and 50 µg/ml subtilisin applied simultaneously (Fig. 4C,D), suggesting that the underlying ion channels are not gated by tip links, kinocilial links or ankle links (Goodyear and Richardson, 2003; Goodyear et al., 2005). Figure 4C provides an example of an observation that was also made in many other hair cells, namely that a smaller current of a similar “peaky” shape at negative potentials (the size of which could vary markedly during a recording in a given cell) was also seen in response to positive stimuli, so that a sine-wave stimulus resulted in current responses with double the stimulus frequency ("2f" responses). In a few cells the anomalous response (in terms of absence of a resting MET current, size and kinetics: see further on) was to positive stimuli, just as for the normal MET current (this was observed in three homozygous mutant OHCs: one Cdhl23−/− and two Myo7a−/−).

Interestingly, the MET currents recorded in response to step stimuli still showed evidence of strong “adaptation,” a decline in current over time, at −84 mV, but not +86 mV, in the presence of 5 mM BAPTA in both the Cdhl23−/− and Myo7a−/− homozygotes. Figure 5 shows recordings from a homozygous Cdhl23−/− OHC at hyperpolarized and depolarized membrane potentials. The step responses in this cell correspond to the responses observed to sine-wave stimuli, with the largest currents elicited by negative force steps (Fig. 5A,B) and smaller currents with similar kinetics seen in response to positive force steps (Fig. 5C,D). For both stimulus directions and at both potentials, ∼50 nm of bundle displacement is required before a current occurs (Fig. 5E). The decline in current is as reported before for these mutants in normal extracellular Ca2+ (Kros et al., 2002; Alagramam et al., 2011), and shows that in these cells this phenomenon does not depend on influx of extracellular Ca2+. The onset kinetics of the currents were also slowed compared with normal MET currents, and as noted before for these mutants this feature is particularly evident at depolarized membrane potentials (Fig. 5B,D).

The anomalous MET currents likely represent a population of MET channels that are gated by mechanisms other than by tip links. To further substantiate this idea we sought to irreversibly damage the hair bundles of normal OHCs by mechanical overstimulation. After doing so we were able to record the anomalous currents following a time interval of between 7 s and 3 min (Fig. 6A–C). Such responses were often unstable during a recording, varying in size and in the degree to which a component responding to positive stimuli was present. Figure 6D–F shows an example of a cell in which a 2f response was recorded transiently before the currents occurred only in response to negative force stimuli.

The anomalous currents could thus be elicited under a variety of conditions: after cleaving the tip links with BAPTA, in mutants lacking functional tip links, and following mechanical overstimulation and destruction of the hair bundle. The responses were usually, but not always, largest for negative force stimuli and a resting MET current could not be observed at hyperpolarized membrane potentials. The threshold for eliciting the anomalous currents at −84 mV was some −50 to −150 nm. When applying force steps, currents developed more slowly than normal and an unusually strong decline was observed at negative potentials (Kros et al., 2002; Alagramam et al., 2011).
Development of normal and anomalous OHC MET currents

We hypothesized that these unusual currents might represent a population of MET channels in an abnormal position that is not gated via the tip-link complex and wondered whether the development of these currents matched that of the normal MET currents. In the apical coil before P2, MET currents are absent or very small in rat (Waguespack et al., 2007). We followed the development of normal (Fig. 7A,B) and anomalous (Fig. 7C–F) MET currents in apical OHCs between E18.5 and P10. The former developed as reported previously with little or no current until P2, and half-maximum size was reached at P3.6 (Fig. 7B). Following application of 3 μM FM1-43 for 10 s (Gale et al., 2001) dye loading was not observed in OHCs (and IHCs) before P2 in the apical coil (data not shown), indicating the absence of channels with a significant probability of being open when the bundles are in their resting position. Anomalous MET currents were already observed at E18.5 in apical-coil OHCs of wild-type mice, sized from -30 to -500 pA at -84 mV (Fig. 7C), and could be elicited by strong mechanical stimuli at a time when normal MET currents were still absent or very small (a few tens of pA). In a sample of 19 E18.5 to P1 OHCs from wild-type CD-1 mice only three cells had small normal MET currents (<30 pA at -84 mV), whereas strong stimulation elicited the anomalous MET current in 11 cells. One cell first showed a small normal MET current (-35 pA) followed by a small anomalous current (-30 pA) during strong stimulation. By P3–P4 the vast majority of cells expressed normal MET currents (Fig. 7A,B). A set of recordings in which 214 P3–P4 OHCs situated in the apical and basal coils of CD-1 mice were stimulated with 40 V sinusoids to elicit saturating MET currents was scrutinized for anomalous MET currents. Four of these cells only showed anomalous MET currents, whereas normal and anomalous MET currents were recorded simultaneously in another six cells. Overall, anomalous MET currents were thus observed rarely under these conditions, in <5% of OHCs. Following BAPTA treatment to sever the tip links however, we reliably observed large anomalous currents in wild-type OHCs between P3 and P10 (Fig. 7C,D). Current size was very variable but there was no obvious trend with age. This also applied to anomalous MET currents recorded between P1 and P5 from homozygous mutant mice with hair-bundle defects (Kros et al., 2002; Alagramam et al., 2011).

Permeation properties of the anomalous channels differ from those of normal MET channels

The rare recordings of OHCs in which anomalous and normal MET currents coexisted offered an opportunity to compare their properties under identical conditions. In these cases the anoma-
lous currents can be distinguished from the normal MET currents by their peaky appearance due to their rapid and declining appearance at negative membrane potentials, and by their occurrence during the negative phase of the stimulus waveform (Fig. 9A, B). The normal MET current is more rounded in appearance at all potentials, and is partially activated at rest as indicated by the reduction of the holding current during the first negative half-cycle of the sine wave. When plotting the current–voltage relation of the anomalous and normal MET currents in 4 OHCs which were studied in 1.3 mM extracellular Ca$^{2+}$ (Fig. 9C), we noticed that the normal double-rectification (inward and outward), which has been attributed to a voltage-dependent block by divalent cations (Kros et al., 1992), is not a feature of the anomalous current. The normal current could be fitted with the single-energy-barrier model used before (Kros et al., 1992; Gale et al., 2001; Marcotti et al., 2005), but the anomalous currents could not and were better fitted with a straight line. This finding suggests that the ion channel that underlies the anomalous MET current may have ion permeation characteristics that are different from those of the normal MET channels.

To gain a more detailed impression of the properties of the conducting pore of the ion channels that carry the anomalous MET currents, we examined block of the currents by the amino-glycoside antibiotic DHS. This has been studied in detail for normal MET currents, for which the half-blocking concentration at a membrane potential of $-84$ mV was reported as 7.0 $\mu$M (Marcotti et al., 2005). Superfusion of 100 $\mu$M DHS caused only a modest reduction of the MET currents recorded from OHCs after their tip links had been broken by exposure to 5 mM BAPTA with, as for the normal MET currents, the block being relieved at depolarized membrane potentials (Fig. 10A–C). Considerably higher concentrations of DHS, in the millimolar range, were required to achieve a nearly complete block at hyperpolarized membrane potentials, which was again largely removed upon depolarization (Fig. 10D–F; an OHC superfused with 3 mM DHS). Such concentrations, when applied to control OHCs, caused a rapid deterioration of the cells precluding successful patch-clamp recordings. In control experiments run in parallel with normal, non-BAPTA-treated OHCs, 10 $\mu$M DHS caused a substantial block at hyperpolarized membrane potentials that was relieved at positive, and also extreme negative, potentials (Fig. 10G–I). Such relief of block at extreme negative membrane potentials, the signature for permeant block of the MET channel (Gale et al., 2001; Marcotti et al., 2005; van Netten and Kros, 2007), was not observed for the anomalous MET currents (Fig. 10C,F). DHS block was found to be reversible upon washout of the drug, both for anomalous and normal MET currents. The dose–responses curves for DHS block of the anomalous MET currents were shifted by two orders of magnitude compared with those of the normal MET currents (Marcotti et al., 2005), with half-blocking concentrations ($K_{D}$) of 894 $\mu$M and 7.0 $\mu$M at $-84$ mV, respectively. At $-164$ mV the $K_{D}$ became 11.4 $\mu$M for the normal MET currents due to partial release of block, but such a shift did not occur for the anomalous MET currents ($K_{D}$, 832 $\mu$M).

Another aminoglycoside antibiotic, neomycin, also caused little block of the anomalous MET currents of OHCs of homoygous Myo7a$^{d1l}$ and Myo7a$^{621658}$ mutant mice (1 $\mu$M blocked 32.89 $\pm$ 0.04%, $n = 3$, of the anomalous current at $-84$ mV). Just as for DHS, block was considerably reduced compared with normal MET currents in three control OHCs from P6 CD-1 mice (100 $\mu$M blocked 72.1 $\pm$ 2.4%, $n = 3$).

**Discussion**

**Anomalous MET currents of OHCs and IHCs**

Our observations show that MET channels in mammalian auditory hair cells can be gated when tip links are removed pharmacologically or are destroyed by repeated mechanical overstimulation. The currents measured under these conditions are of abnormal polarity in that they are elicited predominantly when the bundle is moved away from the kinocilium, and their onset kinetics are slower than normal. At negative membrane potentials, there is little or no resting transducer current and responses to a step stimulus decline rapidly, even for large stimuli. This decline is unlike classic MET current adaptation in that it does not depend on influx of extracellular Ca$^{2+}$ through the MET channels (Crawford et al., 1989, 1991; Beurg et al., 2010; but see Peng et al., 2013). We could also elicit currents with these characteristics before the time at which normal MET currents are reported to appear during development (Waguespack et al., 2007; Lelli et al., 2009; Kim and Fettiplace, 2013). Finally, such currents are observed in OHCs of mutant mice that lack either of the two tip-link proteins, CDH23 or PCDH15 (Alagramam et al., 2011), or the tip-link-associated motor protein MYO7A (Kros et al., 2002; Grati and Kachar, 2011). Our present finding that the anomalous MET currents in the OHCs of homoygous Myo7a$^{d1l}$ mutant mice are resistant to BAPTA treatment (Fig 4C) prompts a reinterpretation of the findings reported by Kros et al. (2002). We must now conclude that the MET currents in these mutants are not gated by means of tip links and are predominantly of opposite polarity relative to normal MET currents. The resistance of the currents to BAPTA and subtilisin, as well as their presence even in bundles that have been mechanically destroyed, would also seem to exclude gating by any of the other links that are situated in the hair bundle at this stage of development (Goodyear et al., 2005).
Location of the ion channels underlying the anomalous MET currents

Through what kind of ion channels could the anomalous MET current flow? Possibilities include the normal MET channels, but in an abnormal position or configuration so that they are not gated by tip links, or other mechanosensitive channels, or altogether different ion channels that are not related to the MET channel. A stretch-sensitive current (I_{metL}) of a few tens of pA that flows through non-selective channels (passing cations as well as anions) has been reported in response to mechanical stimulation by a glass probe or a fluid jet of the basolateral membrane of mature guinea pig OHCs, where it modulates electromotility (Rybalchenko and Santos-Sacchi, 2003). There is also some evidence for stretch-sensitive non-selective cation channels and potassium channels in the basolateral membrane of mature guinea pig OHCs, again with similarly small whole-cell currents (Ding et al., 1991; Iwasa et al., 1991). However, given that the anomalous MET currents are much larger, need strong mechanical stimulation close to the apical surface of the hair cell, occur well before the onset of electromotility around P8 in mouse OHCs (Marcotti and Kros, 1999) and are observed in both OHCs and IHCs, it is unlikely that they are due to these channels in the basolateral membrane.

Our finding of MET currents with inverse response polarity in mammalian hair cells at the earliest stage of their development (Fig. 7) bears comparison with recent observations on developing zebrafish hair cells (Kindt et al., 2012). Although MET currents were not measured in that study, Ca\(^{2+}\) entry into the hair cells was observed to occur early on in development when the bundles were moved away from the kinocilium. The Ca\(^{2+}\) responses in the early immature zebrafish hair cells were different from our OHC currents in that they were eliminated by BAPTA treatment and were absent in mutant zebrafish that lacked either CDH23 or PCDH15. The inverted responses in zebrafish hair cells were also not found in zebrafish mutants that lack normal kinocilia, which prompted Kindt et al. (2012) to propose that the kinocilial links, which are composed of PCDH15 and CDH23 like tip links but are of opposite orientation (Ahmed et al., 2006; Goodyear et al., 2010; Lelli et al., 2010), gate these currents. Given the resistance to treatment by BAPTA (and subtilisin) and the presence of the anomalous currents in mutants lacking tip-link proteins, we consider a location of the channels at the kinocilium, or an association with any other links, unlikely for mouse OHCs. The absence of any clear trend in the size of the anomalous MET currents with age up to P10, the oldest age tested (Fig. 7C,D), also argues against this current being a transient, develop-
mental phenomenon in mammalian hair cells. The abnormal kinetic properties of the currents, their instability, and the lack of a resting current point to an inefficient and variable engagement of the channels. Because the currents can be recorded once the hair bundle has been stripped of it links or mechanically destroyed, we propose that the ion channels underlying the anomalous current are located at the basal insertion points of the stereocilia or elsewhere along the apical membrane overlying the cuticular plate and are activated by the membrane shearing relative to the underlying cytoskeleton. Ion channels with reversal potentials close to zero and single-channel conductances similar to those of the MET channels have been found on the apical membrane of OHCs (Frolenkov et al., 2004).

**Molecular nature of the ion channels underlying the anomalous MET currents**

Our finding that the anomalous MET currents of OHCs are two orders of magnitude less sensitive to block by the aminoglycoside antibiotics DHS and neomycin suggests that the pore properties of the underlying ion channels are different. DHS is a permeant blocker of the normal MET channels (Marcotti et al., 2005), but the lack of release of block at extreme negative potentials suggests that this is not the case for the anomalous MET channels. The current–voltage curves of the anomalous MET currents are linear, unlike the normal MET currents which show a double rectification attributed to permeant block by the divalent cations Ca\(^{2+}\) and Mg\(^{2+}\) (Kros et al., 1992). Very similar anomalous MET currents have recently been reported for OHCs with mutations in the genes encoding the TMC1 and TMC2 proteins (Kim et al., 2013), which are both candidate for being subunits of the MET channel (Kawashima et al., 2011; Barr-Gillespie and Nicolson, 2013; Kim and Fettiplace, 2013; Pan et al., 2013). In homozygous TMC1 and TMC2 mutant mice the anomalous channels had a reduced Ca\(^{2+}\) permeability compared with the normal MET channels, but in contrast to the findings of the current study a relatively normal block by DHS (Kim et al., 2013). The reasons for the latter discrepancy are unclear and merit further investigation. The high Ca\(^{2+}\) permeability, large conductance and high-affinity permeant block by aminoglycoside antibiotics of normal MET channels have been attributed to the presence of a vestibule lined with negative charges at the extracellular side of the channel that concentrates

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**Figure 8.** Anomalous MET currents in a normal P1 OHC in response to force steps. A, B, MET currents recorded at membrane potentials of \(-84\) mV and \(+86\) mV from a P1 apical-coil OHC from a CD-1 mouse. Transducer currents (inward at \(-84\) mV; outward at \(+86\) mV) are elicited by force steps in the negative direction (negative driver voltage). At \(+86\) mV some MET channels are open at rest and are closed by positive force steps. Current declines strongly at \(-84\) mV and slow onset kinetics are particularly evident at \(+86\) mV (activation time constants \(-2.5\) ms). C, Absolute size of peak MET current as a function of driver voltage to the fluid jet at \(-84\) mV (closed circles) and \(+86\) mV (open circles), for a range of force steps.

**Figure 9.** Normal and anomalous MET currents coexisting in the same cells. A, B, MET currents recorded from CD-1 OHCs in 1.3 mM extracellular Ca\(^{2+}\). Membrane potential stepped between \(-164\) and \(+76\) mV in 20 mV increments from a holding potential of \(-84\) mV; for clarity only responses to every other voltage step are shown. Driver voltage (45 Hz sinusoid, 40 V amplitude) plotted above the current traces. In both cells, anomalous MET currents (arrow) are superimposed on the normal MET current (arrowhead), which has a component activated at rest that deactivates during the negative phase of the sine-wave stimulus. C, Current–voltage curves of the normal and coexisting anomalous MET currents of four CD-1 OHCs (P3–P4) in 1.3 mM Ca\(^{2+}\), including those of A and B. Currents are normalized to values at \(+96\) mV: normal MET current \(I_{\text{M, norm}}\) 737 ± 249 pA; anomalous MET current \(I_{\text{M, anom}}\) 415 ± 151 pA. Fits are as follows: normal MET current \(I = 183 \text{pA, } V_r = -10.1 \text{mV, } V_i = 38.5 \text{mV, and } \gamma = 0.46\); anomalous MET current straight line with slope of 3.76 pA/mV.
and funnels cations into the pore (Beurg et al., 2006; van Netten and Kros, 2007; Pan et al., 2012). The low-affinity block by extracellular DHS of the anomalous MET currents is comparable to that of the block of the normal MET currents by intracellular DHS at positive potentials (Marcotti et al., 2005; van Netten and Kros, 2007). The anomalous MET channels therefore seem to lack the vestibule of the normal MET channels. Although it is possible that the anomalous MET current is unrelated to the
normal MET current, a more compelling possibility is that these anomalous currents represent a ‘reserve’ population of MET channel precursors that are not usually evident in the presence of normal MET currents and can be observed when normal mechanotransduction is not yet established or cannot occur due to damaged or absent tip links. These hypothetical precursors have a permeation pore but lack a vestibule, which could conceivably be formed by TMC1 and TMC2 once the channels are in their normal position at the tips of the stereocilia. When a normal MET current is present the anomalous channels are only rarely observed (Fig. 9), suggesting that they are normally suppressed by an as yet unidentified mechanism.

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