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Position-dependent patterning of spontaneous action potentials in immature cochlear inner hair cells

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Abstract

Spontaneous action potential activity is crucial for mammalian sensory system development. In the auditory system, patterned firing activity has been observed in immature spiral ganglion cells and brain-stem neurons and is likely to depend on cochlear inner hair cell (IHC) action potentials. It remains uncertain whether spiking activity is intrinsic to developing IHCs and whether it shows patterning. We found that action potentials are intrinsically generated by immature IHCs of altricial rodents and that apical IHCs exhibit bursting activity as opposed to more sustained firing in basal cells. We show that the efferent neurotransmitter ACh, by fine-tuning the IHC's resting membrane potential ($V_{\rm m}$), is crucial for the bursting pattern in apical cells. Endogenous extracellular ATP also contributes to the $V_{\rm m}$ of apical and basal IHCs by activating SK2 channels. We hypothesize that the difference in firing pattern along the cochlea instructs the tonotopic differentiation of IHCs and auditory pathway.

The origin and development of neural circuits, including those of sensory systems such as the cochlea and the retina, depend on intrinsic genetic programmes that are coordinated by specific microRNAs¹-². While the initial wiring of the sensory circuits involves axon guidance molecules³.⁴, their refinement towards mature functional characteristics is hypothesized to be driven by intrinsically-generated (i.e. experience-independent) electrical activity, which occurs during a critical period of development in immature sensory organs⁵-⁷. Early spontaneous action potential activity has been shown to regulate a variety of cellular responses⁸ including the remodelling of synaptic connections ⁷,⁹ and ion-channel expression¹⁰. As well as promoting the survival of immature sensory cells and newly formed connections within the organ, spontaneous action potential activity is believed to serve a more instructive role in sensory refinement⁹. Differences in pattern, which depend on the cell's position within the immature organ, might help establish ordered neural maps of the sensory environment before the onset of experience-driven activity.

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In the mammalian cochlea, the primary sensory receptor inner hair cells (IHCs), fire spontaneous action potentials^{11_15} before the onset of hearing, which in various altricial rodents occurs around postnatal day 12 (P12). Any differences in the pattern of action potential activity as a function of position along the immature cochlea might instruct the functional differentiation of the IHCs themselves¹⁶, the biophysical properties of which are known to vary along the cochlea^{17_19}, as well as the possible refinement of tonotopic maps in the auditory pathway²⁰. In this study we compared action potential activity in apical and basal IHCs and found position-dependent differences in its patterning along the cochlea. Moreover, we found that while ATP is involved in maintaining $V_{\rm m}$ near the potential at which the Ca²⁺ current begins to activate, the efferent neurotransmitter ACh is likely to be responsible for the burst-like action potential pattern observed specifically in apical IHCs.

Results

In the mammalian cochlea spontaneous spiking activity in immature IHCs has been observed during the first postnatal week^{12,14}. Although action potentials (APs) are mainly due to the interplay between a Ca²⁺ and a delayed rectifier K⁺ current^{14,21}, their frequency has been shown to be modulated by other currents that are transiently expressed in immature IHCs: the Na⁺ current¹⁴ and a small conductance Ca²⁺ activated K⁺ current SK2 (I_{SK2})^{16,22}. Action potential activity is also regulated by the efferent neurotransmitter acetylcholine (ACh)²³. Recently, it has been suggested that early postnatal IHCs have a significantly more hyperpolarized $V_m^{24,25}$ than previously reported using whole-cell^{12,14} or perforated-patch¹⁶ recordings. As a consequence, it was proposed that IHCs are not intrinsically spontaneously active but that instead APs are only triggered by waves of ATP released from supporting cells in Kölliker's organ^{24,25}. These waves were shown to lead to ATP-dependent activation of purinergic receptors in IHCs resulting in a depolarizing inward cation current.

Action potentials occur spontaneously in immature cells

We have investigated the origin of action potential activity in IHCs during the first postnatal week in mice, rats and gerbils at 35-37°C and in perilymph-like extracellular solution (1.3 mM Ca²⁺ and 5.8 mM K⁺) using whole-cell current clamp (Fig. 1a) or cell-attached voltage clamp (Fig. 1b). The cell-attached recording technique has the advantage that it preserves the intracellular milieu, including endogenous Ca^{2+} buffers. Therefore, IHC V_m and the frequency and pattern of spontaneous APs should, as closely as possible, represent that in vivo. During cell-attached recording, APs manifest as biphasic capacitative currents (Fig. 1b) and could be recorded in nearly all IHCs tested within the first postnatal week. Cellattached spikes resulted from Ca^{2+} -dependent APs since they were abolished by the application of a Ca²⁺-free solution (Fig. 1c). Furthermore, the frequency of spikes increased when IHCs were depolarized by extracellular application of a high- K^+ solution (Fig. 1d). Under all conditions tested, the patterns of cell-attached and whole-cell spikes appeared qualitatively the same (Fig. 1). The cell-attached mode allowed spontaneous action potential activity to be recorded for much longer durations (our longest recording was one hour) than whole-cell recording (less than twenty minutes, probably due to Ca^{2+} current rundown), allowing for a more comprehensive analysis. These recordings also represent the first demonstration of long-lasting action potential activity in IHCs, indicative of well-preserved cells with a stable $V_{\rm m}$. The action potential activity occurred at the cell's $V_{\rm m}$ and was not accompanied by any obvious waves of depolarization (Fig. 1, Supplementary Fig. 1) that could be linked to ATP release from Kölliker's organ²⁴,²⁵. This confirms our previous studies using both whole-cell¹⁴,²¹ and perforated patch recordings¹⁶. Similar responses were observed whether the patch pipette was advanced from the modiolar or strial side of the cochlea, the latter ensuring Kölliker's organ remained entirely intact²⁴. However, we noticed that the action potential width in our recordings (half width 3.8 ms in both perforated patch

and whole-cell recordings: Fig. 2a, b) was about 3-9 times shorter than that recently reported²⁵, suggesting differences in the currents underlying APs between the studies.

To establish the basis for the above mentioned differences, we compared recordings made at room temperature with an intracellular solution containing 10 mM EGTA (as recently used to measure IHC voltage responses 24_{26}), to those with our standard conditions of body temperature and 1 mM EGTA. Under the room temperature and 10 mM EGTA condition, we found that ISK2 was absent or significantly reduced compared to that measured in 1 mM EGTA (Fig. 2c, d: one-way ANOVA, Tukey test: P<0.001) or using perforated patch (Fig. 2d: P < 0.01, overall significance P < 0.0001). This indicates that 10 mM EGTA buffers most of the Ca²⁺ entering the IHC before it reaches the SK2 channels. The total outward K⁺ current from IHCs recorded with 10 mM EGTA was similar to that obtained in SK2 knockout mice (perforated patch) and when I_{SK2} was abolished using a Ca²⁺-free extracellular solution²² (whole-cell) (Fig. 2d). In the absence of I_{SK2} the action potential width was significantly increased compared to that measured in perforated patch recordings or using 1 mM intracellular EGTA (Fig. 2b: one-way ANOVA: P < 0.0001). Since I_{SK2} is essential for robust action potential repolarization, its absence affects the ability of an IHC to sustain spiking activity¹⁶,²². At room temperature and with 10 mM EGTA we also found that the $V_{\rm m}$ of apical-coil IHCs was significantly more hyperpolarized (-70.4 ± 1.5 mV, n =11, P3-P4) than that of similar cells recorded at body temperature and using 1 mM EGTA $(-58.5 \pm 0.4 \text{ mV}, n = 44, \text{P2-P5}, P < 0.0001)$, preventing them from firing spontaneous APs. Moreover, we observed discrete bursts of action potential activity associated with brief periods of cell depolarization (9.4 \pm 2.1 mV from five P3 IHCs: Fig. 2e), which were similar to those recently reported under analogous conditions²⁴-²⁶. Therefore, to avoid adversely affecting the nature and kinetics of the currents underlying spontaneous action potential activity, we performed all our whole-cell recordings as previously described^{14,16,21} (35-37°C; 1 mM intracellular EGTA; in perilymph-like extracellular solution²⁷: 1.3 mM Ca^{2+} and 5.8 mM K⁺).

The pattern of spontaneous APs varies along the cochlea

To assess possible position-dependent differences in spontaneous action potential activity, we performed cell-attached recordings from IHCs in the apical and basal regions of the immature rat cochlea (Fig. 3a). The firing pattern of apical IHCs appeared organized in bursts (rat: Fig. 3a, b; gerbil: Supplementary Fig. 2a-c) that were less evident in recordings from basal IHCs (rat: Fig. 3a, c). A consequence of the more burst-like pattern in apical rat IHCs was that the mean but not the median spike frequency (mean 0.60 ± 0.09 Hz, median 2.00 ± 0.14 Hz, n = 17) was significantly lower than in basal cells (mean 1.23 ± 0.13 Hz, median 1.80 \pm 0.13 Hz, n = 15, P < 0.001). In the interspike interval (ISI) histograms (Fig. 3d), the relatively quiet intervals in between the bursts are responsible for the pronounced 'tail' at large ISIs for apical IHCs. We used the coefficient of variation as a quantitative measure of regularity in spontaneous spike firing. The coefficient of variation for a random Poisson process is 1; values <1 indicate more regular activity whereas bursting leads to values >1²⁸, ²⁹. The coefficient of variation in apical IHCs (2.70 \pm 0.24, P2-P5, n = 17) was significantly larger (P < 0.0001) than in basal cells (1.16 ± 0.08, P2-P5, n = 15) (Fig. 3e). We have also observed similar apical-to-basal differences in the frequency and pattern of action potential activity in mouse IHCs (Supplementary Fig. 1), suggesting that they are a robust and general feature of mammalian cochlear development. During the first postnatal week, the coefficient of variation remained relatively constant in apical and basal rat IHCs (Fig. 3f) and apical gerbil IHCs (Supplementary Fig. 2), indicating that the difference in spike pattern is not due to the development of basal cells being about one-to-two days ahead of apical cells³⁰. Gerbils have an extended low-frequency hearing range in the adult (down to ~ 200 Hz), compared to both mice and rats ($\sim 2-3$ kHz). The nevertheless similar

coefficient of variation and frequency in apical IHCs of all three rodent species indicate that the spike pattern relates to the cell's position along the length of the cochlea, rather than to the characteristic frequency of sound to which adult IHCs respond best.

The $V_{\rm m}$ of mouse apical IHCs (-58.5 ± 0.4 mV, n = 44, P2-P5) was found to be significantly more hyperpolarized than that of basal cells (-55.5 ± 0.5 mV, n = 23, P3-P5, P < 0.0001: see also Fig. 7c:"Ctrl"). When small current injections (14 ± 2 pA, n = 4) were applied to apical-coil IHCs, causing them to depolarize by about 3 mV and thus mimicking the $V_{\rm m}$ of basal cells, their firing activity became more regular (Supplementary Fig. 3). This indicates that the mechanisms underlying the different $V_{\rm m}$ in apical and basal IHCs are crucial determinants of their different firing patterns.

Concentration-dependent roles for ATP in spiking activity

During the first (but not the second) postnatal week the $V_{\rm m}$ of immature IHCs (Fig. 4a) is positive to the potential at which the Ca²⁺ current becomes activated (-65 mV^{14} , 31), so at this stage there is no requirement for ATP-induced depolarization to reach the action potential threshold. Because we did not observe any slow waves of depolarization of $V_{\rm m}$ associated with spontaneous APs, we sought to distinguish whether endogenous ATP was simply not present or whether the IHCs did not depolarize in response to ATP. When superfused with 100 μ M ATP mouse IHCs depolarized (Fig. 4b) to a significantly larger extent in basal than in apical IHCs (P < 0.0005: Fig. 4c). ATP reversibly increased the frequency and reduced the height of APs (apical IHCs) or even abolished them (basal IHCs). In the presence of ATP the cell-attached spike amplitude of apical IHCs (7.8 ± 0.3 pA, n =6, Fig. 4d) was reduced by $35 \pm 14\%$ (P<0.001) when compared to that of control spikes $(12.2 \pm 0.8 \text{ pA}, n = 6)$, presumably as a consequence of IHC depolarization. The spike amplitude histogram shows a clear segregation between spikes recorded before and during the application of ATP (Fig. 4e: obtained from the recording in Fig. 4d). Spikes recorded without extracellularly applied ATP had stable amplitudes (Fig. 4f). This shows that action potential bursts are unlikely to result from waves of ATP-induced IHC depolarization.

Having established that IHCs respond to externally applied ATP under our conditions, we sought to investigate the presence of endogenous ATP by applying TNP-ATP, a specific antagonist for purinergic P2X receptors³². The expectation was that IHCs would hyperpolarize if endogenous ATP was present due to the block of inward cation current flowing through P2X receptors. Surprisingly, we found that 10 µM TNP-ATP caused apical (Fig. 5a) and basal (Fig. 5b) IHCs to depolarize instead, reversibly increasing the action potential frequency. Basal IHCs were depolarized more (P < 0.0005) than apical cells (Fig. 5c). A qualitatively similar effect was observed when TNP-ATP was used at lower concentrations (1 nM - 1 µM: Supplementary Fig. 4). The depolarizing action of TNP-ATP was confirmed by the increased frequency and reduced height of spikes in cell-attached recordings (Fig. 5d). Apical-coil IHCs were also similarly depolarized (7.8 \pm 4.1 mV, n = 3) when superfused with the more general purinergic receptor blocker PPADS (50 μ M). When recordings were performed using SK2-knockout mice or experimental conditions that prevented the activation of the SK2 current, 10 µM TNP-ATP had no or very little effect on $V_{\rm m}$ (Fig. 5e, f). This indicates that Ca²⁺ influx into IHCs induced by endogenous ATP is able to activate nearby SK2 channels located at or near the synaptic region (Fig. 5g), thus sustaining a repolarizing outward K⁺ current. P2X receptors and SK2 channels showed very close co-localization of about 13 nm (apical IHCs: Fig. 5e) to 18 nm (basal IHCs: Fig. 5f), as calculated from the BAPTA concentrations of 10 mM and 5 mM, respectively, required to uncouple them²²,³³. When experiments were performed at room temperature and using 10 mM EGTA in the intracellular solution (Fig. 5e), TNP-ATP induced IHC depolarization was reduced or absent, indicating that under these conditions the SK2 channels are no longer

coupled to the P2X receptors. It is worth noting that in the presence of 1 mM EGTA the size of the SK2 current, when elicited by depolarization (Fig. 2c), was not affected by the extracellular application of either TNP-ATP (n = 5) or PPADS (n = 3) (data not shown).

To explain the depolarizing effect of TNP-ATP (Fig. 5) we hypothesized that endogenous ATP (i.e. ATP at very low concentrations) should hyperpolarize IHCs. When P4 apical IHCs were superfused with micromolar concentrations of ATP ($3 \mu M - 100 \mu M$) all 9 mouse IHCs tested were depolarized (Fig. 6a, e). ATP in the 30 nM - 300 nM range caused a small hyperpolarization averaging -2.1 mV (Fig. 6b, e) in 6 out of 22 IHCs. In the remaining 16 IHCs, ATP caused a small depolarization (n = 9, Fig. 6c, e) or had no effect (n= 7). Superfusion of 3 nM - 10 nM ATP (Fig. 6d, e) had a hyperpolarizing effect (-4.1 ± 0.7) mV, range -0.7 mV to -7 mV) in 9 out of 11 IHCs investigated, with two cells showing no response. This hyperpolarization was prevented when TNP-ATP (10 μ M) was superfused together with a low concentration of ATP (3 nM) in six P4 apical IHCs (data not shown). The variable responses at intermediate ATP concentrations are likely to be due to differences in the levels of endogenous ATP in the nanomolar range already present around the IHCs. These findings confirm our observations that endogenous ATP modulates the action potential activity intrinsic to IHCs by activating P2X receptors coupled to SK2 channels located at the synaptic region. These receptors are likely to be different to those responding to high concentrations of ATP (Fig. 4) since in the presence of both TNP-ATP and apamin (to block the P2X receptors and SK2 channels responsible for IHC hyperpolarization) the depolarization induced by 100 µM ATP in apical IHCs (data not shown) remained similar to that recorded when ATP was superfused alone (Fig. 4c).

Role of α9α10 nicotinic ACh receptors in spiking activity

ACh is the main efferent neurotransmitter in the mammalian cochlea that exerts an inhibitory effect on IHC responses before hearing onset²³. Efferent fibres synapse directly with IHCs mainly during pre-hearing stages of development³⁴. Efferent inhibition of IHCs is achieved by Ca²⁺ influx through heteromeric a9a10 nAChRs^{35,36} activating the hyperpolarizing SK2 current^{22,23,37}.

We investigated whether endogenously released ACh had a direct role in modulating intrinsic action potential activity by superfusing spontaneously spiking IHCs with the specific a9a10 nAChR blocker strychnine³⁸. Strychnine (10 µM) caused a small IHC depolarization, sufficient to change the bursting firing pattern characteristic of apical cells into a more sustained activity (Fig. 7a, b). In the presence of strychnine (1 µM - 10 µM), the $V_{\rm m}$ of apical IHCs (control -58.5 ± 0.4 mV, n = 44; strychnine: -54.8 ± 0.7 mV, n = 6) was more depolarized and similar to that of basal cells (control: -55.5 ± 0.5 mV, n = 23; strychnine: -54.1 ± 0.7 mV, n = 4, Fig. 7c). During strychnine application the coefficient of variation (Fig. 7d) and mean frequency (Fig. 7e) of action potential activity in apical IHCs were similar to those of basal cells. Moreover, the $V_{\rm m}$ and action potential properties of apical and basal IHCs in the presence of strychnine did not differ significantly from those of control basal cells (one-way ANOVA, Fig. 7c-e). Cell-attached spike recordings further confirmed that strychnine increased the frequency and decreased the coefficient of variation of action potentials in apical IHCs (Fig. 7f, g) and that the efferent neurotransmitter ACh inhibited IHC firing activity (Fig. 7h).

The more pronounced effect of strychnine on apical-coil IHCs could in principle be due to larger currents induced by ACh or the presence of more endogenously released ACh than in the basal coil. In the presence of 5 mM BAPTA, which uncouples most of the SK2 channels from the ACh receptors²², the inward current induced by 30 μ M ACh measured at -84 mV³⁹ was similar between apical (-63 ± 7 pA, *n* = 5) and basal (-92 ± 24 pA, *n* = 6) P3

IHCs, suggesting a comparable number of ACh receptors. Moreover, $\alpha 9\alpha 10$ nAChRs are likely to be similarly colocalized to SK2 channels in both cochlear regions since 30 µM ACh caused an equivalent degree of hyperpolarization in both apical (14.0 ± 0.1 mV, *n* = 3) and basal (13.0 ± 0.7 mV, *n* = 3) P5 IHCs. The conclusion is that endogenously released ACh is likely to be much more prevalent in the apical coil.

We verified that the effect of strychnine was due to ACh released by the efferent system by attempting to block its release by either preventing action potential activity in the efferent terminals using the Na⁺ channel blocker TTX or by indirectly exhausting exocytosis from the efferent terminals by inhibiting endocytosis of ACh-filled vesicles with dynasore⁴⁰. In the continuous presence of TTX or dynasore (Supplementary Fig. 5), 10 μ M strychnine had very little or no depolarizing effect on all apical IHCs tested (TTX: 0.3 ± 0.1 mV, *n* = 11, P4; dynasore: 1.1 ± 0.6 mV, *n* = 4, P4), when compared to that obtained in control conditions (Fig. 7c: 3.7 mV). The coefficient of variation and mean frequency of action potential activity in apical IHCs in the presence of TTX or dynasore (coefficient of variation: 0.7 ± 0.1; frequency: 5.0 ± 0.4 Hz, *n* = 8, Supplementary Fig. 5) were similar to those of control basal cells (Fig. 7d, e, respectively). These findings indicate that ACh released from the efferent terminals is essential for setting the bursting firing pattern in apical IHCs by tuning their *V*_m to a value more hyperpolarized than that of basal cells.

Discussion

We show that the action potential activity during the first postnatal week in immature IHCs of altricial rodents is intrinsically generated by the cells themselves, but modulated in different ways by inhibitory effects due to ACh and ATP released near the IHCs. The frequency and pattern of this spontaneous activity differs between apical and basal regions of the cochlea, with apical IHCs showing a bursting pattern and basal IHCs a more regular pattern with a higher mean frequency. Patterned spontaneous action potentials have been observed before the onset of hearing in mammalian spiral ganglion cells²⁸,⁴¹ and auditory brain stem neurons in the chick, where position dependence of their frequency was noted⁴². The difference in pattern of spiking activity we have shown along the immature cochlea would be appropriate for guiding tonotopically the functional differentiation of IHCs¹⁷.¹⁹, the specific re-organization of synaptic connections⁴³ and possibly the refinement of sensory maps along the auditory pathway before the onset of experience-driven activity²⁰,⁴².

An important question is: what determines the patterning of IHC action potentials and the differences in pattern, bursting-like versus more sustained firing, along the cochlea? IHC action potential activity is known to be modulated by the inhibitory efferent neurotransmitter ACh^{23} ,⁴⁴ and ATP released from supporting cells²⁴. Our experiments with the ACh receptor blocker strychnine indicate that spontaneous release of ACh from efferent nerve fibres²³, via the indirect activation of SK2 channels, keeps apical IHCs more hyperpolarized than basal cells. The physiological consequence of this is that, by fine-tuning the V_m of apical IHCs, ACh has a specific role in determining the bursting-like action potential pattern in these cells (Supplementary Fig. 6). Although we have demonstrated that endogenous ACh originates from the efferent fibres, and that this release is more pronounced in the apical coil of the cochlea, it is not known whether ACh is itself released in bursts or how this pathway normally operates in the intact animal. Previous *in vivo* studies suggest that input to IHCs from the cholinergic efferent feedback system could be involved in generating patterned activity in auditory nerve afferents⁴⁵,⁴⁶.

ATP also affects IHC action potential activity during the first postnatal week, but in a more complex way than previously thought. Higher concentrations of ATP (in the order of μ M) depolarize the IHCs as previously described²⁴, presumably via P2X₂ receptors localized at

the cell's cuticular plate⁴⁷ (Supplementary Fig. 6). However, we have shown that endogenous ATP around IHCs has, similar to ACh, a hyperpolarizing effect on their $V_{\rm m}$ by activating IHC P2X receptors that indirectly open co-localized SK2 channels present at or near the cell's presynaptic site (Supplementary Fig. 6). P2X₇ and P2X₃ are known to be transiently expressed in developing IHCs⁴⁷. The low nanomolar sensitivity of the P2X receptors to TNP-ATP indicates that P2X₃ is the most likely candidate to mediate the hyperpolarizing ATP responses in immature cells³². P2X₃ receptors show an apical to basal expression gradient along the mouse cochlea (low to high)⁴⁸. This could explain the more pronounced effects of ATP in basal IHCs, which have more sustained action potential firing and may thus require additional assistance in robustly repolarizing their membrane potential. Recordings from guinea-pig IHCs show that both hyperpolarizing and depolarizing effects of ATP also occur in the mature cochlea⁴⁹, indicative of other, non-developmental, roles.

Tonotopic differences in IHC activity during the first postnatal week could be important for refining the auditory neural circuits in the brainstem, most of which is completed during this time window²⁰. From the second postnatal week intrinsic IHC activity disappears due to a progressive hyperpolarization of $V_{\rm m}$. At this time, bursts of action potentials elicited by waves of ATP released from supporting cells²⁴ could be important for coordinating activity in adjacent IHCs. Developmental changes in the mechanisms underlying patterned activity have been found in the retina, spinal cord and hippocampus⁷, to which the cochlea can now probably be added. We have shown that there are numerous influences on IHC action potential activity that together exert tight control over the IHCs' intrinsically generated firing pattern as a function of position along the immature cochlea. The need for such elaborate control over the $V_{\rm m}$ of IHCs points to the crucial importance of their firing activity during cochlear development.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Appendix

Methods

Electrophysiology

Apical and basal coil inner hair cells (IHCs) from the mouse (n = 338), rat (n = 43) and gerbil (n = 27) were studied in acutely dissected organs of Corti from postnatal day 2 (P2) to P11, where the day of birth is P0. Animals were killed by cervical dislocation in accordance with UK Home Office regulations. Cochleae were dissected as previously described²¹ 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, 2 sodium pyruvate, amino acids and vitamins (pH 7.5; osmolality ~308 mmol kg⁻¹). Superfusion of IHCs with a Ca²⁺-free solution, 15 mM K⁺, 3 nM-100 μ M ATP, 1 nM-10 μ M TNP-ATP, 50 μ M PPADS, 300 nM TTX (Tocris, UK), 10-100 μ M ACh, 1-10 μ M strychnine, 100 μ M dynasore (Sigma) was performed with a multi-barrelled pipette positioned close to the patched cells. TTX or dynasore were applied for at least 10-20 minutes before using either strychnine or ACh (Supplementary Fig. 5). The spike amplitude was normally distributed in control

experiments and best fitted with a single Gaussian (Fig. 4e, f). In experiments in which ATP was extracellularly applied, the spike amplitude distribution was best fitted with two Gaussian functions.

The pipette solution used for cell-attached recordings contained (in mM): 140 NaCl, 5.8 KCl, 1.3 CaCl₂, 0.9 MgCl₂, 0.7 NaH₂PO₄, 5.6 D-glucose, 10 Hepes-NaOH (pH 7.5). The whole-cell pipette solution contained (mM): 131 KCl, 3 MgCl₂, 1 EGTA-KOH, 5 Na₂ATP, 5 Hepes-KOH, 10 sodium phosphocreatine (pH 7.3). In some cases 1 mM EGTA was replaced by 10 mM EGTA or different BAPTA concentrations (5 mM and 10mM) and KCl was adjusted to keep the osmolality constant. The following KCH₃SO₃-based intracellular solution (in mM: 134 KCH₃SO₃, 1 MgCl₂, 10 EGTA-KOH, 2 Na₂ATP, 20 Hepes-KOH, 0.2 Na₂GTP, pH 7.3) was used for recording from apical-coil IHCs in order to test whether the different experimental conditions previously used elsewhere^{24_26} to record action potential activity from IHCs (mainly room temperature and 10 mM intracellular EGTA) affected the cells' responses. Data from perforated-patch experiments (Fig. 2) were obtained by analyzing recordings from IHCs previously obtained for Refs ¹⁶, ²². Patch pipettes were made from Soda glass capillaries (Harvard Apparatus Ltd) and coated with surf wax (SexWax, Mr Zoggs) to minimise the fast transient due to the patch pipette capacitance. Unless otherwise stated, experiments were performed close to body temperature (35-37°C) and with 1.3 mM Ca²⁺ and 5.8 mM K⁺ in the extracellular solution, similar to those of perilymph²⁷, to try and approach normal physiological conditions as best as possible.

Electrophysiological recordings were made using Optopatch (Cairn Research Ltd, UK) or Axopatch 200B (Molecular Devices, USA) amplifiers. Data acquisition was controlled by pClamp software using Digidata 1322A or 1440A boards (Molecular Devices, CA, USA). Recordings were low-pass filtered at 2.5 kHz (8-pole Bessel) and sampled at 5 kHz. Cellattached action potential recordings (seal resistance was 1.80 ± 0.05 GO, n = 149) were filtered at 10 kHz and sampled at 20 kHz. Some were in addition low-pass filtered offline at 500 Hz (8-pole Bessel) using Clampfit. Data analysis was performed using Origin software (OriginLab, MA, USA). The Mini Analysis Program (Synaptosoft Inc. NJ, USA) was used to detect biphasic spike events, to calculate their frequency and to analyze interspike intervals (ISI) and amplitudes. The action potential frequency was calculated as the reciprocal of the mean ISI for each cell and an indication of the spread of ISI values about the mean was obtained by calculating the coefficient of variation, equal to the standard deviation divided by the mean. Whole-cell membrane potentials (V) were corrected for the liquid junction potential of -4 mV (KCl-based intracellular) or -10 mV (KCH₃SO₃-based intracellular), measured between electrode and bath solutions.

Statistical analysis

Statistical comparisons of means were made by Student's two-tailed *t*-test or, for multiple comparisons, one-way ANOVA followed by the Tukey test. Means are quoted \pm s.e.m. and P < 0.05 was used as the criterion for statistical significance.

Immuno-fluorescence labelling

Cochleae from immature mice were used to prepare cryosections for immuno-fluorescence microscopy and processed as previously described⁵⁰. Briefly, cochleae were dissected and fixed for 2 h with 2% paraformaldehyde (w/v), embedded and cryosectioned at a thickness of 10 μ m. Sections were mounted with Vectashield medium with DAPI (Vector Laboratories). Antibodies to SK2 channels (rabbit, Sigma, diluted 1:50) and the IHC marker otoferlin (mouse, Abcam, diluted 1:100) were used. Primary antibodies were detected with Cy3-conjugated (Jackson ImmunoResearch Laboratories) or Alexa Fluor 488–conjugated secondary antibodies (Molecular Probes). Sections were viewed using an Olympus BX61

microscope equipped with motorized z axis and epifluorescence illumination. Images were acquired using a CCD camera and analyzed with cellSense Dimension software (OSIS). To display SK2 channel distribution, cochlea slices were imaged over a distance of several μ m to cover the IHC width in an image-stack along the z axis (z stack) followed by three-dimensional deconvolution (cellSense Dimensions, OSIS). Fig. 5g shows composite images, which represent the maximum intensity projection over all layers of the z stack.

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Figure 1. Cell-attached spikes result from spontaneous APs

(**a** and **b**), Spontaneous APs recorded with whole-cell current clamp and cell-attached voltage clamp, respectively. Right panels show an action potential on an expanded time-scale. (**c** and **d**), Whole-cell (top panels) and cell-attached (bottom panels) recordings of spontaneous APs in mouse apical-coil IHCs during the application of a Ca²⁺-free solution containing 0.5 mM EGTA (**c**), a condition that abolishes this activity¹⁴. During the perfusion of a solution containing 15 mM K⁺ (**d**) IHCs depolarized causing the frequency of spikes to increase while their amplitude declined. In this and the following figures, the duration of substance application is indicated by the horizontal line above the recording.

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Figure 2. Spontaneous action potentials in IHCs require physiological recording conditions (a), Spontaneous APs recorded from mouse IHCs maintained at 35-37°C and using 1 mM intracellular EGTA^{14,16,21} (black traces) or at room temperature using 10 mM EGTA^{24,26} (grey). (b), Width of the APs measured at 50% between the peak and the after-hyperpolarization. Action potential width in 10 mM EGTA (n = 9, P3-P4) was similar to that measured in IHCs from SK2 knockout mice (SK2^{-/-}: n = 9, P4) but larger than that obtained in perforated patch and whole-cell with 1 mM EGTA (n = 5, P3-P7). Number above the symbols indicates the APs analyzed. In this and the following figures, error bars indicates s.e.m. (c), Currents in voltage clamp obtained using 1 mM EGTA before and

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during the superfusion of a Ca²⁺-free solution, a condition that abolishes $I_{SK2}^{16,22}$ (arrowhead). Recordings were obtained by applying 4 s voltage steps to -10 mV from a holding potential of either -74 mV or -84 mV^{22} . In the presence of 10 mM EGTA (Ref: ²⁴-²⁶), I_{SK2} was absent or largely reduced. (**d**), Size of the total K⁺ current (I_K :delayed rectifier and I_{SK2}) from IHCs under different recording conditions. Dashed horizontal line indicates that I_{SK2} is normally superimposed onto I_K . Asterisks indicate significance compared to control. (**e**), At room temperature and using 10 mM EGTA single or bursts of APs were elicited by slow periods of membrane depolarization.

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Figure 3. Patterning of spontaneous spiking activity in apical and basal IHCs (a), Cell-attached recordings of spontaneous spikes from an apical and basal rat IHC (120 s out of: apical 31 min; basal 6 min). (b and c), ISIs as a function of time from the same apical and basal IHCs shown in a (mean frequency: apical 0.26 Hz, basal 2.40 Hz; median frequency: 2.32 Hz, 2.68 Hz; coefficient of variation: 2.8, 0.65). The bursting nature of spiking activity in apical IHCs (b) is indicated by periodic increases in spiking rate in between silent/reduced activity. In basal IHCs (c) spike discharge was more continuous and ISIs were less variable than in apical cells. (d), ISI histograms (bin size 50 ms) from the same IHCs shown in a. Inset shows the full scale x-axis for the apical IHC. The mode was 0.35 ms for both the apical and basal IHCs. (e), Coefficient of variation from each IHC against their firing rate (apical, closed symbols: basal, open symbols; averages in grey). (f), Average coefficient of variation as a function of postnatal age. Symbols are as in e. IHCs (P2-P5) used for analysis in e and f were: 17 apical (recording time: 3.6 hrs, 5726 spikes) and 15 basal (1.2 hrs, 5504 spikes).

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Figure 4. Modulation of action potential activity by superfusing ATP

(a), $V_{\rm m}$ of immature apical IHCs (closed symbols) measured with whole-cell recording. Open triangles indicate the membrane potential threshold required for triggering APs (n = 68, P2-P11). (b), ATP increased the action potential frequency in early postnatal apical (upper panel) and basal (lower panel) IHCs, using whole-cell recordings. (c), Shift in $V_{\rm m}$ when apical and basal IHCs were superfused with 100 μ M ATP. (d), ATP increased spike frequency and decreased their height in apical IHCs using cell-attached recordings. (e), Histograms (bin size 0.5 pA) showing the spike amplitude before and during the superfusion of ATP from the same IHC shown in d (recording time: 514 s, 377 spikes). (f), Spike amplitude histogram from an apical IHC in the absence of exogenous ATP (recording time: 869 s, 748 spikes).



Figure 5. Role of endogenous ATP in spontaneous spiking activity in mouse IHCs (a and b), Spontaneous APs recorded with whole-cell current clamp from apical and basal IHCs, respectively, during superfusion of TNP-ATP. Note the increased frequency and decreased height of APs during the application of TNP-ATP, with basal IHCs being more affected than apical cells. (c), Depolarizing shift in $V_{\rm m}$ of apical and basal IHCs during the superfusion of TNP-ATP. (d), Spontaneous cell-attached spikes recorded from a mouse IHC in the presence of TNP-ATP. (e), Comparison of $V_{\rm m}$ shift when apical P3-P5 IHCs were superfused with TNP-ATP alone or together with the SK2 channel blocker apamin (both with 1 mM intracellular EGTA), in SK2 knockout mice (SK2^{-/-}) or when the intracellular Ca²⁺ buffers BAPTA (5-10 mM) or EGTA (10 mM) were used. The experiments with 10 mM EGTA were performed at room temperature. (f), $V_{\rm m}$ shift when basal IHCs were perfused with TNP-ATP alone or in the presence of 5 mM intracellular BAPTA. (g), Apical immature IHC immunostained for SK2 channels (red) and the IHC marker otoferlin (green). Nuclei were stained with DAPI (blue). Scale bar: 10 μ M

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Figure 6. Dual action of ATP on spontaneous spiking activity in mouse IHCs

(a), Spontaneous action potentials recorded with whole-cell current clamp from an apical IHC. Extracellular application of 3 μ M ATP caused IHC depolarization. (**b** and **c**), High nanomolar concentrations of ATP had mixed effects on IHC $V_{\rm m}$, with some cells showing a small hyperpolarization (**b**) and others exhibiting a small depolarization (**c**: slightly increased action potential frequency). (**d**), Low nanomolar concentrations of ATP (3 nM) caused IHC hyperpolarization. Note that in some cases following ATP application a small depolarizing rebound occurred before the IHC returned to a stable $V_{\rm m}$. (**e**), $V_{\rm m}$ shift in response to different ATP concentrations.

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Figure 7. Effect of endogenous ACh on IHC spontaneous spiking activity

(a), Spontaneous whole-cell action potentials from an apical P4 IHC with the superfusion of 10 μ M strychnine, a specific a9a10 nAChR blocker. Strychnine caused a small IHC depolarization, which increased the action potential frequency. (b), ISIs as a function of time from the same apical IHC shown in **a**. Note the periodic increases in spiking rate with strychnine in between silent/reduced activity. (**c**-e), $V_{\rm m}$ (**c**), coefficient of variation (**d**) and frequency (**e**) of action potentials in control conditions and with 1 μ M or 10 μ M strychnine measured from P2-P5 apical and basal IHCs. For all three panels, significant differences were only observed between apical control IHCs and the other conditions (overall significance from one-way ANOVA: $V_{\rm m} P < 0.0001$; coefficient of variation and frequency P < 0.005). (**f**), Cell-attached spikes showing increased firing activity in the presence of 1 μ M strychnine. (**g**), As shown for whole-cell recordings (**d**), in the presence of strychnine the coefficient of variation (CV) of cell-attached spike ISIs in apical IHCs was significantly lower than control apical cells (P < 0.05) but similar to that of control basal cells. (**h**), Cell-attached spikes are abolished during the application of the inhibitory efferent neurotransmitter ACh.