

Tonotopic Variation in the Calcium Dependence of Neurotransmitter Release and Vesicle Pool Replenishment at Mammalian Auditory Ribbon Synapses

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The mammalian cochlea is specialized to recognize and process complex auditory signals with remarkable acuity and temporal precision over a wide frequency range. The quality of the information relayed to the auditory afferent fibers mainly depends on the transfer characteristics of inner hair cell (IHC) ribbon synapses. To investigate the biophysical properties of the synaptic machinery, we measured changes in membrane capacitance (ΔC_m) in low-frequency (apical region, ~ 300 Hz) and high-frequency (basal, ~ 30 kHz) gerbil IHCs maintained in near physiological conditions (1.3 mM extracellular Ca^{2+} and body temperature). With maturation, the Ca^{2+} efficiency of exocytosis improved in both apical and basal IHCs and was more pronounced in the latter. Prehearing IHCs showed a similar Ca^{2+} cooperativity of exocytosis despite the smaller ΔC_m in apical cells. After maturation, ΔC_m in high-frequency IHCs increased linearly with the Ca^{2+} current, whereas, somewhat surprisingly, the relationship was significantly more nonlinear in low-frequency cells. This tonotopic difference seemed to be correlated with ribbon synapse morphology (spherical in apical and ellipsoid in basal IHCs) but not with the expression level of the proposed Ca^{2+} sensor otoferlin or the spatial coupling between Ca^{2+} channels and active zones. Repetitive stimulation of adult IHCs showed that vesicle pool refilling could become rate limiting for vesicle release, with high-frequency IHCs able to sustain greater release rates. Together, our findings provide the first evidence for a tonotopic difference in the properties of the synaptic machinery in mammalian IHCs, which could be essential for fine-tuning their receptor characteristics during sound stimulation.

Key words: hair cell; ribbon synapse; exocytosis; development; cochlea; calcium current

Introduction

The exquisite temporal acuity of the mammalian cochlea depends on neurotransmitter release at inner hair cell (IHC) ribbon synapses onto auditory afferent fibers (Fuchs, 2005). Ribbon synapses are specialized structures capable of tethering numerous synaptic vesicles at the release sites of IHCs and photoreceptors (Sterling and Matthews, 2005), allowing them to facilitate high rates of sustained synaptic transmission and coordinated release of vesicles [mammals (Glowatzki and Fuchs, 2002; Goutman and Glowatzki, 2007; Neef et al., 2007); bullfrog (Keen and Hudspeth, 2006)]. Therefore, the ability to relay information with high-fidelity to the central auditory pathway is primarily determined by the transfer characteristics at IHC ribbon synapses. This is dependent on vesicle fusion triggered by Ca^{2+} entry through

L-type ($Ca_v1.3$) Ca^{2+} channels (Platzer et al., 2000) clustered at the presynaptic active zones (Roberts et al., 1990; Tucker and Fettiplace, 1995). Although mature IHCs respond to sound stimuli with graded receptor potentials (Cody and Russell, 1987), prehearing cells (approximately P12 in most rodents) fire spontaneous Ca^{2+} -dependent action potentials (APs) (Beutner and Moser, 2001; Marcotti et al., 2003a,b). In order for IHC synaptic transmission to be optimally configured for the different receptor characteristics during immature and adult stages of development, their synaptic machinery has to change extensively (Sobkowicz et al., 1982; Moser et al., 2006). Biophysically, ribbon synapses become more Ca^{2+} efficient (Beutner and Moser, 2001; Johnson et al., 2005) and less Ca^{2+} dependent (Johnson et al., 2005) with maturation, possibly allowing adult IHCs to respond best to continuous and graded stimuli and at the same time broaden their dynamic range.

The sensory neuroepithelium of vertebrate auditory organs is tonotopically organized such that the characteristic frequency (CF) of hair cells (the sound frequency at which they respond best) gradually changes with position along the auditory organ. In lower vertebrates, this is intrinsically dictated by a position-dependent variation in the biophysical properties of the hair cells (Fettiplace and Fuchs, 1999). Because the mechanical tuning of the mammalian cochlea is thought to provide the necessary fre-

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quency selectivity and sensitivity, IHCs are as yet not considered to be intrinsically tuned themselves or to show great tonotopic variation in their properties. However, recent studies have shown such variation in some intrinsic IHC biophysical properties [Ca^{2+} -buffering proteins (Hackney et al., 2005); transducer current (Jia et al., 2007); Ca^{2+} current (Johnson and Marcotti, 2008)]. Despite the central role for IHC synaptic machinery in auditory transduction, there remains a dearth of information regarding the tonotopicity of its biophysical properties. Therefore, we investigated the kinetics and Ca^{2+} dependence of vesicle release in low- and high-frequency gerbil IHCs *in situ* using near physiological recording conditions. Our main findings indicate the existence of tonotopic differences in the Ca^{2+} dependence of exocytosis and vesicle pool replenishment at IHC ribbon synapses with high-frequency cells being more indefatigable than low-frequency cells. The information presented here provides the first evidence for a position-dependent synaptic specialization, the function of which could be to optimize the responses of these auditory receptors.

Materials and Methods

Electrophysiology. Low- and high-frequency IHCs ($n = 120$) from gerbils were studied in acutely dissected organs of Corti from postnatal day 20 (P20) to P69, at which the day of birth is P0. The average age in apical and basal IHCs was P38 and P42, respectively, which is considered to be mature (Woolf and Ryan, 1984; Müller, 1996). A few prehearing apical and basal IHCs were also investigated ($n = 18$; P5–P7). Gerbils were killed by cervical dislocation in accordance with UK Home Office regulations. Cochleae were dissected in normal extracellular solution (in mM): 135 NaCl, 5.8 KCl, 1.3 CaCl_2 , 0.9 MgCl_2 , 0.7 NaH_2PO_4 , 5.6 D-glucose, 10 HEPES-NaOH. Sodium pyruvate (2 mM), amino acids, and vitamins were added from concentrates (Thermo Fisher Scientific). The pH was adjusted to 7.5 (osmolality ~ 308 mmol kg^{-1}). The dissected cochleae were viewed using an upright microscope (Leica Microsystems) with Nomarski differential interference contrast optics. Adult IHCs were positioned at a frequency range of 250–420 Hz in apical and 20–37 kHz in basal cells (Müller, 1996).

Whole-cell patch-clamp recordings were performed at body temperature (34–37°C) using an Optopatch amplifier (Cairn Research). Patch pipettes (2–3 M Ω) were coated with surfax (Mr. Zogs Sex Wax) and contained the following (in mM): 106 Cs-glutamate, 20 CsCl, 3 MgCl_2 , 1 EGTA-CsOH, 5 Na_2ATP , 0.3 Na_2GTP , 5 HEPES-CsOH, 10 Na_2 -phosphocreatine (pH 7.3; 294 mmol kg^{-1}). In some cases, 1 mM EGTA was replaced by different BAPTA concentrations (0.1, 0.3, 0.6, 1, and 5 mM), and when necessary, CsCl was adjusted to keep the osmolality constant. In a few experiments, perforated patch was used, and the pipette-filling solution contained the following (in mM): 110 K-aspartate, 21 CsCl, 3 MgCl_2 , 5 Na_2ATP , 1 EGTA or 1 BAPTA, 5 HEPES-CsOH, 10 Na_2 -phosphocreatine (pH 7.3; 295 mmol kg^{-1}). The antibiotic amphotericin B (Merck) was dissolved in dry DMSO before dilution in the above intracellular solution to 120 or 240 $\mu\text{g ml}^{-1}$ (Johnson et al., 2007).

Real-time changes in membrane capacitance (ΔC_m) were measured as described previously (Johnson et al., 2002, 2005). Briefly, a 4 kHz sine wave of 13 mV rms was applied to IHCs from -81 mV and was interrupted for the duration of the voltage (V) step. The capacitance signal from the Optopatch was filtered at 250 Hz and sampled at 5 kHz. ΔC_m was measured by averaging the C_m trace over a 200 ms period after the voltage step and subtracting the prepulse baseline. Data were acquired using pClamp software and a Digidata 1320A (Molecular Devices) and analyzed with Origin (OriginLab). Calcium currents were corrected offline for the linear leak conductance (1.7 ± 0.1 nS; $n = 135$; P5–P69). Membrane potentials were corrected for the voltage drop across the series resistance (5.2 ± 0.1 M Ω ; $n = 136$) and a liquid junction potential of -11 mV, measured between electrode and bath solutions. The average voltage-clamp time constant (product of R_s and cell membrane capacitance C_m , 10.2 ± 0.1 pF; $n = 137$) was 54 ± 1 μs ($n = 136$).

Experiments were performed in the presence of 30 mM tetraethylam-

monium and 15 mM 4-AP (Fluka) to block $I_{K,f}$ (Kros et al., 1998) and delayed rectifier K^+ currents [$I_{K,neo}$ and $I_{K,s}$ (Marcotti et al., 2003a)]. Apamin (used for some experiments at 300 nM; Merck) and linopirdine (80–100 μM ; Tocris Bioscience) were also used to block I_{SK} in immature (Marcotti et al., 2004) and $I_{K,n}$ in adult (Marcotti et al., 2003a) IHCs, respectively. The Ca^{2+} dependence of ΔC_m was studied by superfusing a Ca^{2+} -free solution (containing 0.5 mM EGTA) or different Ca^{2+} concentrations (0.6, 1.3, 2.5, 5, and 10 mM). When the concentration of blockers or Ca^{2+} that were added to or removed from the solution was >1 mM, NaCl was adjusted to keep the osmolality constant.

Statistical comparisons of means were made by the two-tailed t test or, for multiple comparisons, one-way ANOVA followed by Tukey's test. Two-way ANOVA, followed by the Bonferroni test, was used to compare data sets from apical and basal IHCs. Means are quoted \pm SEM, and $p < 0.05$ indicates statistical significance.

Immunocytochemistry. For immunofluorescence microscopy, the cochleae from adult gerbils were dissected and processed as described previously (Schug et al., 2006). Otoferlin (mouse monoclonal; Biozol), CtBP2 (rabbit polyclonal; Proteintech Group), and polyclonal anti-Neurofilament 200 (NF200; Sigma-Aldrich) antibodies were used for immunostaining. Primary antibodies were detected with cyanine-3-conjugated (Jackson ImmunoResearch Laboratories) or Alexa Fluor 488-conjugated antibodies (Invitrogen). Sections were embedded with Vectashield mounting medium with 4',6'-diamidino-2-phenylindole dihydrochloride (DAPI) (Vector Laboratories) and viewed using an Olympus AX70 microscope equipped with epifluorescence illumination. Images were acquired using a CCD camera and the imaging system analySIS (Soft Imaging System) and processed with Adobe Photoshop. CtBP2-immunopositive ribbons were counted from apical and midbasal IHCs from six gerbils of three different age groups between P22 and 12 months. The distribution of otoferlin in apical and basal IHCs was determined in at least three animals of a given age (P22, P43, and P120) and done at least in triplicate on each.

Electron microscopy. Apical and basal IHCs from P34 gerbil cochleae were fixed by direct perfusion in 2.5% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.3, for 2 h, washed three times in the above buffer, and then decalcified in 4.13% EDTA, pH 7.3, for 2 d at 4°C. Cochleae were then postfixed (2 h) in buffered 1% OsO_4 , partially dehydrated by an ethanol series up to 70%, at which point the cochleae were incubated in saturated uranyl acetate (70% ethanol) overnight, before completing the ethanol dehydration. The whole intact, decalcified cochlea was embedded in plastic and sectioned in a plane parallel to modiolus. At a near mid-modiolar location, thin sections that included the entire height of the cochlea were taken and mounted on Formvar-coated single slot grids, so that sections through both the apical and the basal coil were included in a single thin section. Serial thin sections of 100 nm thickness of the entire cochlea were then cut as closely as possible parallel to the modiolus and each section mounted on a different grid. All sections were stained with uranyl acetate and lead citrate. The sections at the level close to the modiolus itself were investigated so that the longitudinal sections of basal and apical IHCs examined were in approximately the same orientation to the plane of sectioning. The serial sections were displayed in sequence so that most ribbons could be visualized in their entirety. Basal and apical IHCs from the same sections were examined. Synaptic ribbons were visualized from three apical ($n = 10$) and two basal IHCs ($n = 16$). Ribbon dimensions were taken from those captured in a micrograph and oriented to give an accurate measurement (apical, $n = 6$; basal, $n = 9$).

Results

Whole-cell patch-clamp recordings were used to investigate Ca^{2+} -dependent exocytosis in low-frequency (~ 300 Hz) and high-frequency (~ 30 kHz) gerbil inner hair cells maintained *in situ* in near physiological conditions (body temperature and 1.3 mM extracellular Ca^{2+}). Synaptic vesicle exocytosis was measured as an increase in cell C_m that is interpreted as a sign of neurotransmitter release from presynaptic cells (Parsons et al., 1994; von Gersdorff et al., 1996; Moser and Beutner, 2000).

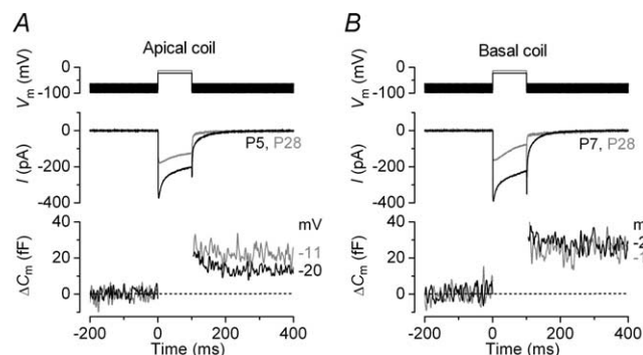


Figure 1. Increased Ca^{2+} efficiency of exocytosis in gerbil IHCs during maturation. **A**, **B**, I_{Ca} (middle) and ΔC_m (bottom) responses recorded from immature (black traces) and adult (gray traces) IHCs positioned in apical and basal regions of the cochlea, respectively. Recordings were obtained in response to 100 ms voltage steps from the holding potential of -81 to -20 mV or -11 mV, where the peak of the I - V curve occurred (Figs. 2B, 3B, bottom). The voltage protocol is shown above the traces. Unless otherwise stated, all recordings were at body temperature and used physiological 1.3 mM extracellular Ca^{2+} .

Position-dependent variation in the Ca^{2+} dependence of exocytosis

Recent studies on mouse apical IHCs (frequency range, ~ 3 – 5 kHz) have shown that the coupling between Ca^{2+} channels and the synaptic machinery becomes more efficient from around the onset of hearing (Beutner and Moser, 2001; Johnson et al., 2005). Therefore, we first investigated whether gerbil IHCs follow a similar developmental pattern. Figure 1A,B shows the maximal Ca^{2+} current (I_{Ca}) and the corresponding ΔC_m recorded in apical and basal IHCs, respectively, at immature and adult stages of development. Although the size of I_{Ca} decreased in both cochlear regions with maturation, the induced ΔC_m was comparable throughout development indicative of an increase in the Ca^{2+} efficiency of exocytosis (qualitatively judged to be $\Delta C_m/I_{\text{Ca}}$). This is suggestive of a common physiological maturation of the IHC synaptic machinery between species. Currently, there is a complete absence of information regarding any position-dependent difference in the properties of neurotransmitter release in IHCs, which if found could greatly improve our understanding of how auditory signals are encoded at the first synapse in the auditory pathway. For these experiments, the gerbil was preferred to the more commonly used mouse because of its extended low-frequency hearing range, which would emphasize any tonotopic differences (gerbil, ~ 0.1 – 60 kHz; mouse, ~ 2 – 100 kHz) (Greenwood, 1990; Müller, 1996).

A direct comparison of I_{Ca} and the corresponding ΔC_m from apical and basal immature IHCs is shown in Figure 2A. Although the size of I_{Ca} was similar between IHCs (apical, -334 ± 23 pA, $n = 9$, P5–P6; basal, -272 ± 25 pA, $n = 7$, P6–P7) (Fig. 2B, bottom) (see also Johnson and Marcotti, 2008), the induced ΔC_m was found to be significantly larger in basal (30 ± 3 fF) than in apical cells (14 ± 1 fF; $p < 0.0001$) (Fig. 2B, top), indicative of a lower Ca^{2+} efficiency of exocytosis in the latter. The synaptic transfer functions (Augustine et al., 1985; Johnson et al., 2005), illustrating the Ca^{2+} dependence of exocytosis (defined as the change in ΔC_m as a function of I_{Ca}) (Eq. 1), are shown in Figure 2C. For this comparison, peak I_{Ca} was preferred to its time integral because, for voltage steps longer than a few tens of milliseconds and at positive membrane potentials, I_{Ca} could be contaminated by the presence of residual unblocked K^+ currents (Johnson et al., 2005, 2007). I_{Ca} inactivation would not affect our interpretations because it has recently been shown to be similar in

apical and basal immature IHCs (Johnson and Marcotti, 2008). Data shown in Figure 2C were approximated using a power function:

$$\Delta C_m = cI_{\text{Ca}}^N, \quad (1)$$

where c is a scaling coefficient, and the exponent N is the power. The power obtained from fitting the data from prehearing IHCs (apical, 3.8 ± 0.3 , $n = 9$; basal, 3.8 ± 0.4 , $n = 7$) indicates that at least four Ca^{2+} ions have to cooperatively bind to the Ca^{2+} sensor to trigger a release event (Dodge and Rahamimoff, 1967; Johnson et al., 2005). This suggests that apical and basal IHCs exhibit a comparable Ca^{2+} dependence of exocytosis despite the smaller ΔC_m in the former. The lower Ca^{2+} efficiency of exocytosis in apical IHCs could be a consequence of fewer vesicles and/or ribbons or more extrasynaptic Ca^{2+} channels compared with basal cells.

Calcium currents and corresponding ΔC_m recordings from low- and high-frequency adult IHCs are shown in Figure 3A. Both the size of the peak I_{Ca} at -11 mV (apical, -151 ± 8 pA, $n = 15$; basal, -145 ± 10 pA, $n = 13$) (Fig. 3B, bottom) and the induced ΔC_m (apical, 20 ± 2 fF; basal, 22 ± 1 fF) (Fig. 3B, top) were similar between apical and basal IHCs. The power value of the synaptic transfer function in basal IHCs (~ 30 kHz) was 1.0 ± 0.1 ($n = 13$) (Fig. 3C,D, bottom), suggesting a linear relationship between Ca^{2+} entry and exocytosis similar to previous findings in apical adult mouse IHCs (~ 3 – 5 kHz) (Brandt et al., 2005; Johnson et al., 2005, 2007) and hair cells of the turtle auditory papilla (~ 100 – 300 Hz) (Schnee et al., 2005). Surprisingly, the power value of adult apical (300 Hz) gerbil IHCs was significantly higher (2.2 ± 0.2 ; $n = 15$; $p < 0.0001$) (Fig. 3C,D, top) than that of basal cells, to some extent resembling that measured in prehearing gerbil cells. It is possible that a proportion of I_{Ca} in IHCs ($>90\%$ carried by $\text{Ca}_v1.3$) (Platzer et al., 2000; Brandt et al., 2003) is driven by Ca^{2+} channels not directly linked to the exocytotic machinery, the distribution of which could differ along the cochlea. However, this is unlikely to affect our results because the theoretical removal of this homogeneous population of channels would possibly compress the transfer functions but not change their power values. A change in the Ca^{2+} dependence of exocytosis would only occur if one or more additional types of Ca^{2+} channel with different voltage-dependent properties and not coupled to vesicle release were expressed in IHCs. Because apical and basal adult gerbil IHCs show a small but significantly different degree of I_{Ca} inactivation (Johnson and Marcotti, 2008), the capacitance synaptic transfer functions (Fig. 3C) were re-plotted using the time integral of I_{Ca} to represent the total Ca^{2+} entry during the 100 ms steps. The power values obtained using I_{Ca} integrals (apical IHCs, 2.5 ± 0.2 ; basal IHCs, 1.2 ± 0.1 ; data not shown) were comparable with those obtained using peak I_{Ca} , suggesting that the amount of Ca^{2+} seen by the synaptic machinery was not significantly affected by I_{Ca} inactivation.

Recent investigations have suggested that the opening of each Ca^{2+} channel allows sufficient Ca^{2+} entry to initiate the release of only very nearby vesicles (Neher, 1998; Brandt et al., 2005). Therefore, the amount of vesicles released summates with the voltage-dependent activation of Ca^{2+} channels. This would give the impression of a linear Ca^{2+} dependence of exocytosis although the machinery may have intrinsically higher-order dependence, which becomes evident at positive membrane potentials (Brandt et al., 2005). Assuming that the intracellular Ca^{2+} concentration at the release site is proportional to the single-channel current (Naraghi and Neher, 1997), the intrinsic Ca^{2+}

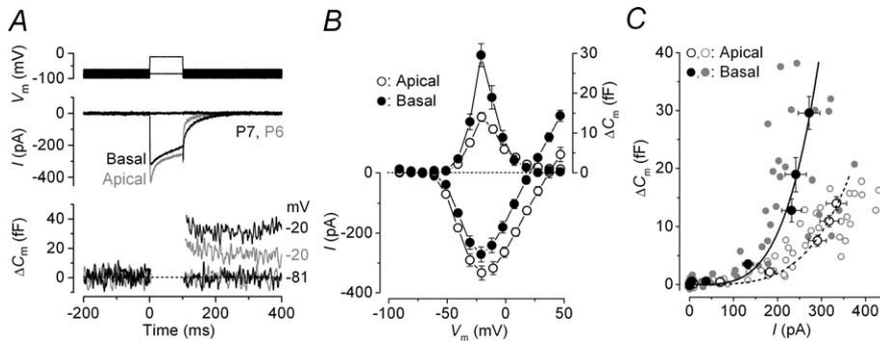


Figure 2. Ca^{2+} currents and ΔC_m in prehearing gerbil IHCs. **A**, I_{Ca} and ΔC_m responses recorded from immature apical (gray traces) and basal (black traces) IHCs. Recordings were in response to 100 ms voltage steps from -81 in 10 mV nominal increments. For clarity, only responses at -81 and -20 mV are shown. **B**, Average peak I - V (bottom) and ΔC_m - V (top) curves from apical (P5–P6; $n = 9$) and basal (P6–P7; $n = 7$) IHCs. **C**, Synaptic transfer functions obtained by plotting average ΔC_m against the corresponding I_{Ca} between -71 mV and near -11 mV from **B** for apical and basal IHCs. Single data points (gray circles) are also shown. The fits are according to Equation 1. Error bars indicate SEM.

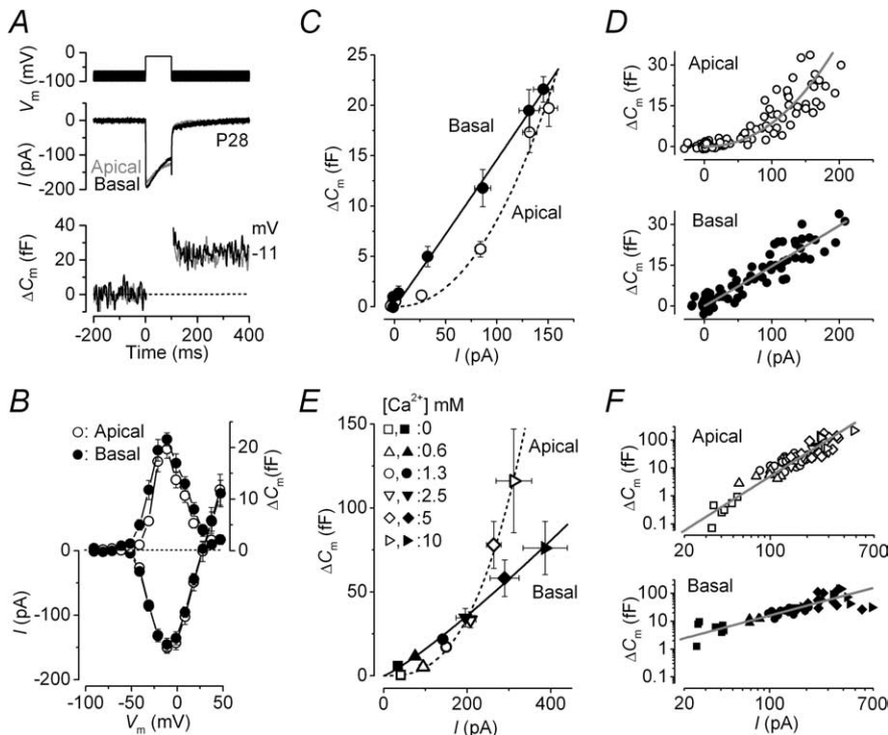


Figure 3. Ca^{2+} dependence of exocytosis in adult gerbil IHCs. **A**, I_{Ca} and ΔC_m from apical (low-frequency, gray) and basal (high-frequency, black) IHCs. Voltage protocol is as in Figure 1. For clarity, only responses at -81 and -11 mV are shown. **B**, Average peak I - V and ΔC_m - V curves in apical (P22–P34; $n = 15$) and basal (P20–P30; $n = 13$) IHCs. **C**, Synaptic transfer functions obtained by plotting average ΔC_m against the corresponding I_{Ca} between -71 mV and near -11 mV from **B** for apical and basal IHCs. The fits are according to Equation 1. **D**, Single data points used to generate averages in **C** from apical and basal IHCs. Fits are as in **C**. **E**, Synaptic transfer functions from apical and basal IHCs showing average responses at the membrane potential at which the peak I_{Ca} from the I - V curve occurred (from approximately -11 to -1 mV) in different Ca^{2+} concentrations. Fits are using Equation 1. **F**, Single data points used to generate averages in **E** from apical and basal IHCs plotted on a double logarithmic scale. Fits are as in **E**. In both **E** and **F**, the number of observations at each Ca^{2+} concentration (0, 0.6, 1.3, 2.5, 5, 10 mM) were as follows: apical 7, 6, 26, 11, 13, 5 (from 26 IHCs); basal 7, 4, 15, 4, 11, 8 (from 16 IHCs). Error bars indicate SEM.

sensitivity of exocytosis in gerbil IHCs could be investigated by recording ΔC_m during the superfusion of different extracellular Ca^{2+} concentrations (Augustine and Charlton, 1986). This would allow ΔC_m to be measured at a membrane potential at which the number of open Ca^{2+} channels is maximal and constant (i.e., at the peak of the I - V) and the single-channel current would be changed only by the extracellular Ca^{2+} concentration. By using this method we avoid possible errors incurred when

evaluating the Ca^{2+} dependence of exocytosis at positive membrane potentials, such as uncertainties about the identity of the current measured over this range (Augustine and Charlton, 1986), the voltage dependence of Ca^{2+} channel gating (Augustine et al., 1985), and the direct effect of membrane potential on the release process (Llinás et al., 1981). Varying the extracellular Ca^{2+} concentration gradually increased the size of both I_{Ca} and ΔC_m in apical and basal IHCs (data not shown). The average I_{Ca} and corresponding ΔC_m values obtained in these different Ca^{2+} concentrations are plotted as synaptic transfer functions in Figure 3E. The power values from fits using Equation 1 for apical and basal adult IHCs were 2.8 ± 0.1 (apical, $n = 26$) and 1.2 ± 0.1 (basal, $n = 16$; significant at $p < 0.0001$), consistent with those obtained using different voltage steps in physiological extracellular Ca^{2+} (1.3 mM) (Fig. 3C,D). Therefore, this suggests that the Ca^{2+} dependence of exocytosis measured in these cells is likely to represent the intrinsic properties of the synaptic machinery. The single data points used for the averages shown in Figure 3E are plotted on a double logarithmic scale (Fig. 3F) to emphasize in more detail responses over the physiological Ca^{2+} range.

Vesicle pool dynamics and ribbon number as a function of IHC frequency position

The rate of neurotransmitter release in adult IHCs was studied by measuring ΔC_m in response to depolarizing voltage steps to -11 mV of varying duration (2 ms to 1.5 s; interstep interval was at least 11 s), which allowed us to investigate the emptying of different synaptic vesicle pool populations (Fig. 4A). Although relatively short stimuli reveal the number of vesicles docked at the active zones [readily releasable pool (RRP)], longer steps induce the release of vesicles from a secondarily releasable pool (SRP) that is located farther away from the Ca^{2+} channels (Parsons et al., 1994; von Gersdorff et al., 1996; Moser and Beutner, 2000; Schnee et al., 2005). Similar to previous findings in mice (body temperature and 1.3 mM Ca^{2+}) (Johnson et al., 2005, 2007), voltage steps of up to ~ 100 ms are likely to recruit mainly the RRP because the increase in ΔC_m could be approximated with a single exponential (Fig. 4B). The size of the isolated RRP was slightly smaller in apical IHCs than in basal cells, as well as the initial release rate (apical, 387 ± 32 ff/s or 10449 ± 877 vesicles/s, $n = 13$; basal, 551 ± 50 ff/s or 14883 ± 1349 vesicles/s, $n = 13$, significant at $p < 0.02$ from fits to individual IHCs), but the SRP in the former seemed to be larger. This behavior appears to reflect the higher-order Ca^{2+} dependence of exocytosis in apical IHCs (Fig. 3E).

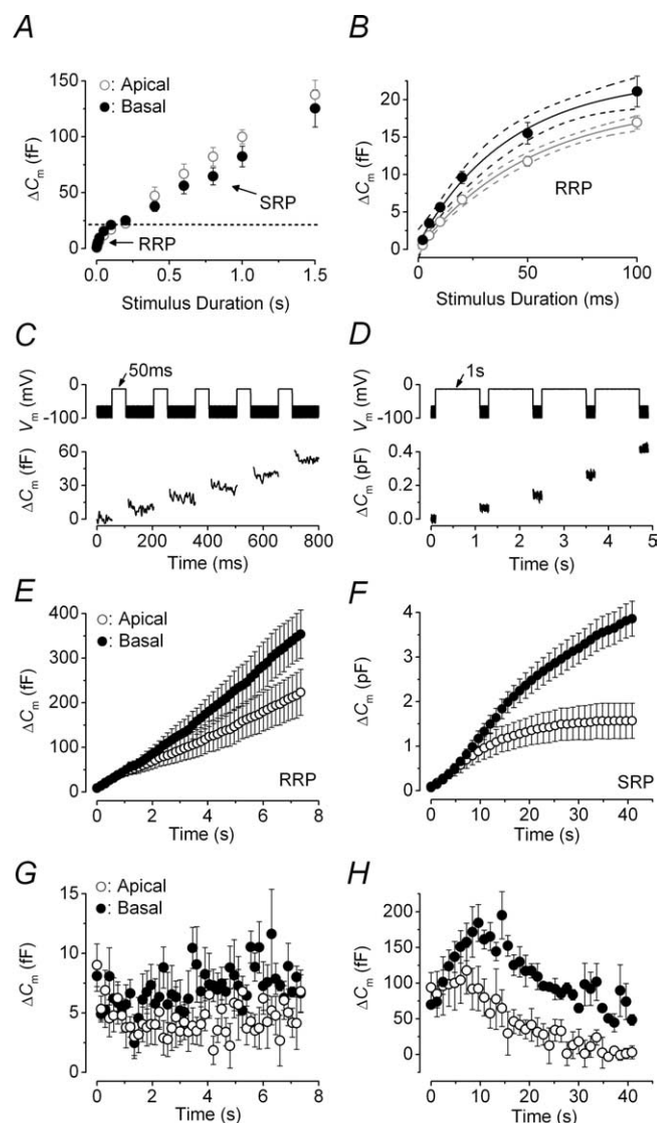


Figure 4. Kinetics of vesicle release and vesicle pool replenishment in adult gerbil IHCs. **A**, Average ΔC_m from 13 apical and 13 basal IHCs (P25–P60) in response to voltage steps from 2 ms to 1.5 s (to approximately -11 mV) showing the RRP and SRP. **B**, RRP (expanded from **A**) approximated with single exponential functions (apical, maximum $\Delta C_m = 20 \pm 1$ fF, $\tau = 52 \pm 9$ ms; basal, maximum $\Delta C_m = 23 \pm 2$ fF, $\tau = 42 \pm 11$ ms). The available RRP consisted of 534 (apical) and 619 (basal) vesicles using a conversion factor of 37 aF/vesicle (Lenzi et al., 1999). Dotted lines represent 95% confidence intervals for the fits. **C, D**, ΔC_m elicited using repetitive voltage steps to -11 mV of 50 ms and 1 s in duration to elicit the RRP and SRP, respectively. The interstep interval was 100 ms in **C** and 200 ms in **D**. For clarity, only the first few steps are shown. The voltage protocol used is shown above the traces. **E, F**, Average cumulative ΔC_m values obtained in response to the 50 ms (50 steps) and 1 s (35 steps) protocol, respectively, from seven apical and three basal IHCs. **G, H**, Individual ΔC_m values from apical and basal IHCs measured after each voltage step from **E** and **F**, respectively. Error bars indicate SEM.

The kinetics of exocytosis measured in this study are unlikely to be affected by slow endocytosis ($\tau = 7.5$ s at room temperature) (Moser and Beutner, 2000) although it is likely to be faster at body temperature.

To calculate the release rate at single ribbon synapses in apical and basal IHCs, we counted them (see Fig. 6A) using an antibody against the presynaptic ribbon component RIBEYE (CtBP2) (Schmitz et al., 2000; Khimich et al., 2005). The total number of labeled ribbons per IHC did not vary significantly along the adult gerbil cochlea (apical, 20.6 ± 0.6 , $n = 20$; basal, 22.4 ± 0.8 , $n = 12$), in agreement with the number of synaptic terminals previ-

ously determined using electron microscopy (Slepecky et al., 2000). Therefore, the vesicle release rate per ribbon was ~ 507 and 664 vesicles/s in apical and basal IHCs, respectively. This indicates that the difference in the total RRP release rate (Fig. 4B) was not attributable to variations in the total number of ribbons per cell or the Ca^{2+} load per synapse because a similar number of Ca^{2+} channels is likely to be present per ribbon.

Vesicle pool depletion could constitute a rate-limiting step in neurotransmitter release. Therefore, we investigated whether vesicle release from either the RRP or SRP was limited by the relative pool-refilling rates using repetitive stimulation. Figure 4, C and D, shows examples of ΔC_m when a train of 50 ms and 1 s steps to -11 mV were applied to investigate the depletion of the RRP and SRP, respectively (Fig. 4A, B). After repeated 50 ms steps, the cumulative ΔC_m showed a near linear increase (Fig. 4E) indicating that the RRP appears to be able to replenish after each step in both apical and basal IHCs. The shallower slope observed in apical IHCs was a consequence of their smaller overall individual ΔC_m (Fig. 4G) (apical, 4.5 ± 0.2 fF; basal, 7.1 ± 0.3 fF; two-way ANOVA; $p < 0.0001$). In contrast to basal IHCs, apical cells exhibited an initial small decline in the individual ΔC_m with stimulus repetition (Fig. 4G), indicating that complete RRP replenishment did in fact not occur within 100 ms. However, when a steady level of release was reached, the machinery was able to replenish this between steps. The first sign of a similar initial ΔC_m decline in high-frequency IHCs was evident using either 50 or 20 ms interstep intervals (data not shown), although for these shorter intervals the recruitment of the SRP became evident, most likely attributable to Ca^{2+} accumulation. These findings suggest that repetitive stimuli could cause some small degree of vesicle pool depletion, which was never complete and was more evident in low-frequency IHCs. The cumulative ΔC_m after long-lasting (1 s) voltage steps showed SRP release saturation that was much more pronounced in apical IHCs (Fig. 4F). The earlier depletion of the SRP in apical IHCs, compared with basal cells, was also evident from the individual ΔC_m (Fig. 4H). Interestingly, repetitive stimulation of the SRP resulted in an initial increase/potential in synaptic transmission before vesicle pool depletion occurred. Qualitatively similar results were also obtained when ΔC_m was normalized to the size of I_{Ca} (data not shown). The observed facilitation could be explained by an initial Ca^{2+} accumulation after long-lasting depolarizations before vesicle depletion occurred.

Functional coupling between Ca^{2+} channels and the RRP

To determine the possible reason behind the different exocytotic Ca^{2+} dependence observed between low- and high-frequency adult gerbil IHCs (Fig. 3), we searched for possible variations in the functional coupling between Ca^{2+} channels and the synaptic machinery. This was done by investigating the morphology of ribbon synapses (Sobkowitz et al., 1982), the colocalization between ribbons and Ca^{2+} channels (Roberts et al., 1990; Tucker and Fettiplace, 1995), and the distribution of the proposed Ca^{2+} -sensor otoferlin (Roux et al., 2006) as a function of IHC CF location. Figure 5, A and B, shows a typical example of the morphology of IHC presynaptic ribbons using transmission electron microscopy. Although all ribbons observed ($n = 16$) in basal IHCs showed an ellipsoid morphology, as described previously in adult mouse IHCs (Sobkowitz et al., 1982; Khimich et al., 2005), those in low-frequency cells were all spherical ($n = 10$). The ribbon width was similar between apical and basal IHCs (spherical ribbons, 114 ± 20 nm, $n = 6$; ellipsoid ribbons, 94 ± 7 nm, $n = 9$; dimensions were only from ribbons in which accurate

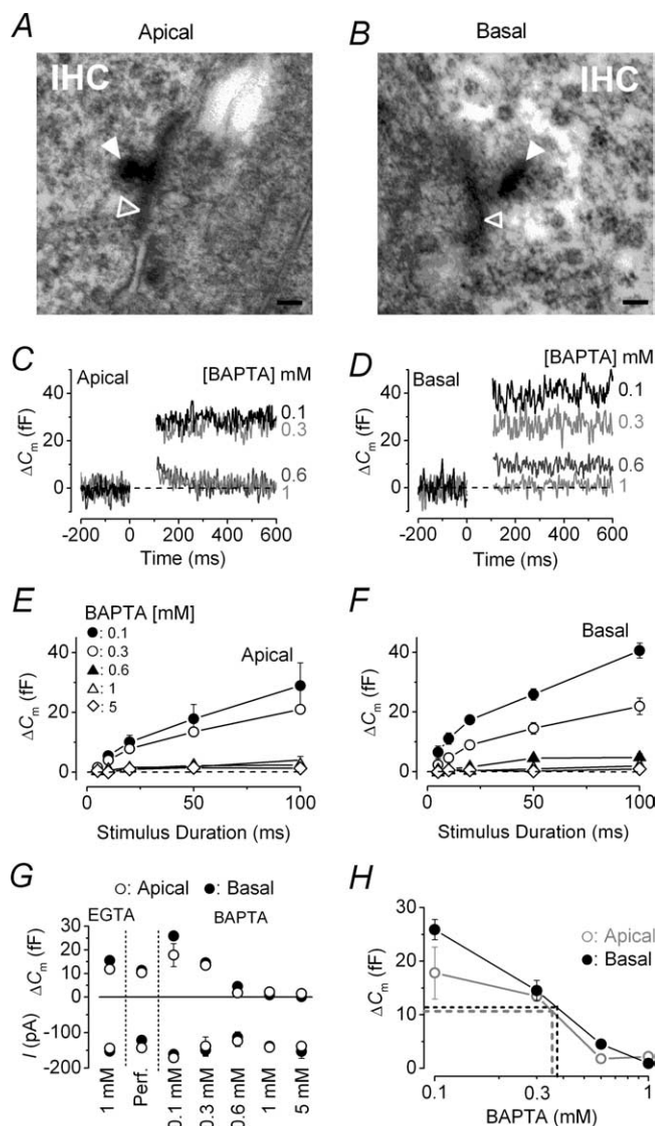


Figure 5. Shape of ribbon synapses and their functional coupling with Ca^{2+} channels. **A, B**, Typical cross-sectional profiles of synaptic ribbons from an apical and a basal IHC, respectively. Note that the ribbon (filled arrowheads) in the apical IHC (**A**) had a round cross-sectional profile, whereas that in the basal cell (**B**) was elongated in shape. Open arrowheads indicate synaptic membrane thickenings. Scale bar, 100 nm. **C, D**, ΔC_m from apical and basal IHCs, respectively, in response to a 100 ms voltage step (to approximately -11 mV) using different intracellular BAPTA concentrations (indicated next to the traces). **E, F**, Average ΔC_m to voltage steps from 5 to 100 ms in apical and basal IHCs. **G**, Average ΔC_m and peak I_{Ca} at 50 ms from data shown in **E** and **F**, including those in EGTA from Figure 4B. Number of cells in EGTA (1 mM), perforated-patch, and different concentrations of BAPTA (0.1, 0.3, 0.6, 1, 5 mM) are as follows: apical, 13, 5, 3, 5, 4, 5, 3; basal, 13, 7, 3, 4, 5, 4, 3. **H**, Estimation of the endogenous buffer concentration. The perforated-patch values of ΔC_m were extrapolated (dotted lines) to those obtained using different BAPTA concentrations (data as in **G**). Error bars indicate SEM.

measurements could be made), but their height was significantly bigger in the latter cells (spherical, 120 ± 19 nm; ellipsoid, 198 ± 10 nm, $p < 0.002$). These dimensions are also comparable with those obtained in other species (Nouvian et al., 2006).

Because the ribbon shape was different between the two CF locations, we investigated whether the colocalization between vesicle release sites and Ca^{2+} channels was affected by measuring ΔC_m in the presence of different intracellular BAPTA concentrations (Fig. 5C–H). BAPTA is a Ca^{2+} chelator with faster binding kinetics than EGTA and therefore capable of buffering Ca^{2+} elevations closer to their source (Neher, 1998). For these experi-

ments, RRP release was studied using a protocol similar to that described in Figure 4B (voltage steps up to 100 ms, RRP) and applied to apical and basal IHCs using different BAPTA concentrations (Fig. 5C,D). The release of the RRP was evident when IHCs were buffered with 0.1 and 0.3 mM BAPTA, but was greatly reduced in both cochlear regions with ≥ 0.6 mM BAPTA (Fig. 5E,F) relative to recordings in 1 mM EGTA (Fig. 5G) ($p < 0.001$), suggesting that a comparable distance is present between Ca^{2+} channels and release sites in both apical and basal cells. Perforated-patch recordings (Fig. 5G) were used to determine whether potential position-dependent differences in the endogenous Ca^{2+} buffering could affect the coupling between Ca^{2+} and exocytosis independent of Ca^{2+} channel position. Figure 5H shows that the concentration of the endogenous mobile Ca^{2+} buffer was ~ 0.38 mM BAPTA in basal IHCs and 0.35 mM BAPTA in apical cells.

Because the relative distance of the Ca^{2+} channels with release sites was comparable in low- and high-frequency IHCs, we investigated whether a difference in the Ca^{2+} dependence of exocytosis could be linked with a tonotopic variation in the expression of otoferlin (Schug et al., 2006), the proposed Ca^{2+} sensor of exocytosis in cochlear hair cells (Roux et al., 2006; Beurg et al., 2008). To determine whether otoferlin was present at IHC active zones, we performed double-staining experiments with either CtBP2 (RIBEYE) (Fig. 6A) or NF200 (Fig. 6B), which stains presynaptic ribbon synapses or afferent fibers, respectively. We found no obvious difference in otoferlin staining along the tonotopic axis of adult gerbil cochleae (P22–P120, qualitative judgment from observations on >200 sections), and although it is clearly involved in exocytosis (Roux et al., 2006; Beurg et al., 2008), it seems unlikely that it accounts for the difference in Ca^{2+} dependence observed in our study. Similarly, the expression of otoferlin in mouse IHCs appears to be constant throughout development (Roux et al., 2006) and again is unlikely to be responsible for the linearization in the Ca^{2+} dependence of exocytosis observed in these cells during maturation (Johnson et al., 2005). Moreover, the apparent absence of otoferlin from the retina (Yasunaga et al., 1999) indicates that it cannot be responsible for the linear Ca^{2+} dependence observed at ribbon synapses of rod photoreceptors (Thoreson et al., 2004). Although otoferlin seemed to be more concentrated below the nucleus of IHCs, it was not exclusively colocalized with either CtBP2-immunopositive ribbons (Fig. 6A, Merge) or NF200 afferents (Fig. 6B). The homogenous distribution of otoferlin throughout IHCs has also been shown in the mouse (Roux et al., 2006) and could indicate a more ubiquitous role for otoferlin in vesicle-membrane trafficking (Yasunaga et al., 1999). Finally, it is also crucial to point out that IHCs from hypothyroid rats did not express otoferlin but still showed exocytosis (Brandt et al., 2007).

Discussion

Exocytosis was investigated at IHC ribbon synapses from the gerbil cochlea using capacitance measurements. Prehearing IHCs and low-frequency (~ 300 Hz) adult cells exhibited a higher-order exocytotic Ca^{2+} dependence than that observed in high-frequency (~ 30 kHz) adult cells. This position-dependent difference in adult IHCs appeared to be correlated with ribbon synapse shape but was not related to the expression level of the Ca^{2+} sensor otoferlin nor the colocalization between Ca^{2+} channels and release sites. Repetitive stimulation of adult IHCs showed that vesicle pool refilling could be rate limiting for the release of the SRP, but not the RRP, with high-frequency IHCs being able to sustain greater release rates. Using the intact gerbil organ of Corti,

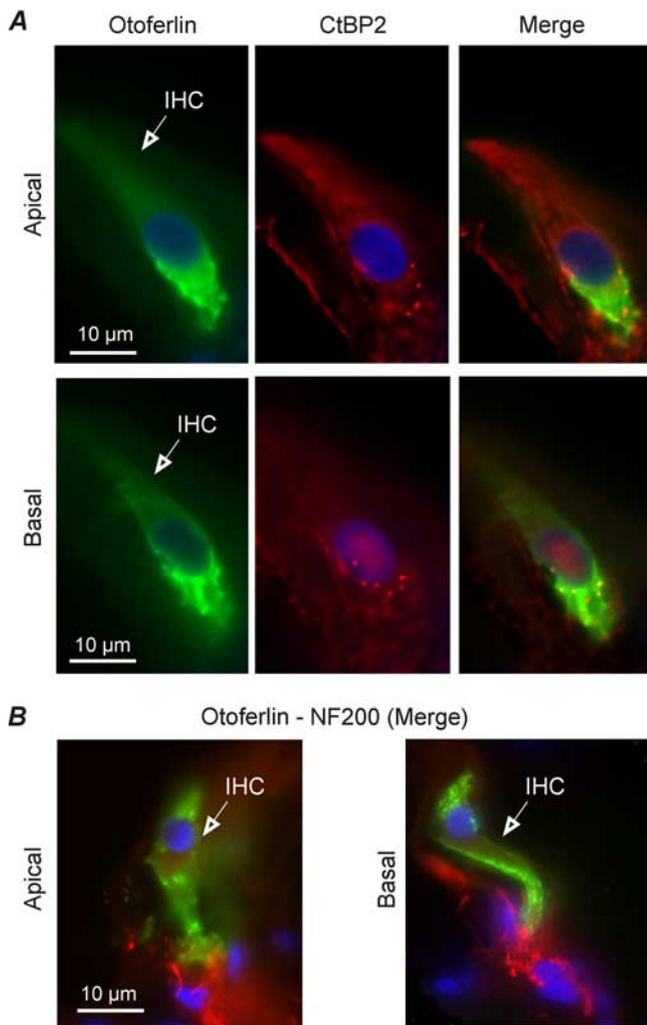


Figure 6. Distribution of otoferlin and ribbon synapses in low- and high-frequency adult gerbil IHCs. **A**, IHCs from apical and midbasal cochlear regions of the adult gerbil (P22) were immunostained for the synaptic vesicle protein otoferlin (green) and presynaptic ribbon component RIBEYE (CtBP2, red). Note that CtBP2 is exclusively localized in the basal pole of IHCs in a dot-like pattern. **B**, IHCs from apical and midbasal cochlear regions (P120) were immunostained for both otoferlin (green) and NF200 (red), which labeled auditory afferent fibers. Nuclei were stained with DAPI (blue). IHCs are indicated by open arrows.

we have provided the first evidence for tonotopic differences in exocytosis at IHC ribbon synapses.

The tonotopicity of gerbil IHC ribbon synapses

In the mouse cochlea, IHC ribbon synapses change from spherical to ellipsoid during maturation (Sobkowicz et al., 1982). We found that although ribbons in high-frequency IHCs of adult gerbils were ellipsoid, those present in low-frequency cells were spherical, suggesting that the latter could retain some immature characteristics. However, the developmental improvement in the Ca^{2+} efficiency of exocytosis in apical and basal IHCs suggests that synaptic maturation did occur in these cells (Beutner and Moser, 2001; Johnson et al., 2005). This leads us to assume that the position-dependent variation in mature ribbon shape is determined by the different functional requirements of low- and high-frequency IHCs and not by the stage of development.

In the mature cochlea, basal-coil IHCs respond to high-frequency sound with depolarizing receptor potentials, initiated by the transducer current at the hair bundle. For high-frequency

tones, cell membrane filtering prevents phase-locking above ~ 3.5 kHz (Palmer and Russell, 1986) so receptor potentials are graded and sustained to represent sound intensity and stimulus envelope. The linear Ca^{2+} dependence ($N \approx 1.0$) and high Ca^{2+} efficiency of the synaptic machinery observed in high-frequency IHCs (~ 30 kHz) ensure that information is accurately encoded at ribbon synapses, as suggested previously in apical mouse IHCs (~ 3 kHz) (Brandt et al., 2005; Johnson et al., 2005) and photoreceptors (Thoreson et al., 2004). In contrast, low-frequency IHCs are phase-locked to sound stimulation such that their receptor potentials are graded to intensity, similar to basal cells, but have an additional phasic component representing the sound frequency (Dallos, 1985). This phasic component, likely to be the dominant component in apical gerbil (~ 300 Hz) but not in apical mouse (~ 3 kHz) IHCs, could be emphasized by the somewhat unexpected finding of a higher-order exocytotic Ca^{2+} dependence in apical cells ($N \approx 2.2$ – 2.8). This view is reinforced by the resemblance of their cooperative nature to that of prehearing IHCs [gerbils (Fig. 2); mice (Johnson et al., 2005)], because AP activity shares some degree of similarity to the phasic nature of low-frequency adult cells. A functional consequence of the higher Ca^{2+} cooperativity of exocytosis in low-frequency IHCs could be the impedance of signaling low-intensity sound stimuli and may contribute to the higher auditory threshold of these cells (Ryan, 1976). Another functional representation of variations in IHC exocytosis could be in the firing characteristics of single auditory afferent fibers that are known to vary tonotopically along the gerbil cochlea, potentially extending the dynamic range of firing rate responses in high-frequency fibers (Ohlemiller and Siegel, 1994). Although some of these differences (spontaneous and sound induced firing rates) could originate from tonotopic variations in IHC exocytosis, it is difficult to correlate these data because afferent activity is driven by single release sites, whereas IHC exocytosis is an integral over the entire cell. Moreover, other factors including postsynaptic efferent modulation and different IHC receptor characteristics could also influence afferent responses.

A compelling observation was the apparent relationship between ribbon morphology and IHC receptor characteristics, with spherical ribbons mainly supporting phasic neurotransmission whereas the ellipsoid shape appears to promote sustained and graded release (Sterling and Matthews, 2005; Moser et al., 2006). However, the possible mechanism involved in linking ribbon shape and function is currently unknown. In contrast to the gerbil, most lower vertebrate auditory hair cells that respond phasically to sound and appear to have mainly spherical ribbons [lizard (Mulroy, 1986); chick (Martinez-Dunst et al., 1997); turtle (Schnee et al., 2005); bullfrog (Keen and Hudspeth, 2006)] show a linear Ca^{2+} dependence of exocytosis (Schnee et al., 2005; Keen and Hudspeth, 2006) that does not vary with CF (Schnee et al., 2005). However, a greater Ca^{2+} cooperativity was evident for membrane potentials negative to -55 mV that elicited small I_{Ca} [see Keen and Hudspeth (2006), their Fig. 5C], probably reflecting the intrinsic properties of the synaptic machinery in these hair cells. The kinetics of exocytosis in turtle hair cells do show tonotopic variation that can be modeled independent from exocytotic Ca^{2+} sensitivity (Schnee et al., 2005). However, our findings suggest that the Ca^{2+} dependence of exocytosis could be more fundamental in determining tonotopic differences in mammalian neurotransmitter release and should be considered when modeling the synaptic vesicle cycle.

Mechanisms controlling Ca^{2+} dependence at IHC synapses

The tonotopic variation in the Ca^{2+} dependence of exocytosis in adult gerbil IHCs is likely to result from a different functional coupling between Ca^{2+} entry and vesicle release, which could be determined by their relative distance (Brandt et al., 2005). In this situation, the Ca^{2+} dependence of the synaptic machinery would be intrinsically high order, regardless of cochlear position, but a closer (nanodomain) coupling to the channels in basal IHCs could give an “apparent” impression of a linear Ca^{2+} dependence over a physiological voltage range. Exocytosis in gerbil IHCs was almost completely prevented using 0.6 mM BAPTA, suggesting that docked vesicles are within 50 nm from Ca^{2+} channels in both cochlear regions (Naraghi and Neher, 1997; Neher, 1998). Therefore, a nanodomain control of exocytosis is likely to be present regardless of CF location, which seems to contradict the above hypothesis. However, we found that the tonotopic difference in the apparent Ca^{2+} dependence of exocytosis (Fig. 3C) seemed to be an intrinsic property of the IHCs synaptic machinery (Fig. 3E), possibly resulting from the differential expression of Ca^{2+} -sensing molecules or promoters of exocytosis (Jahn et al., 2003). Although otoferlin is clearly involved in exocytosis (Roux et al., 2006; Beurg et al., 2008), our findings, together with recently published evidence (see Results), makes it an unlikely candidate for the linearization of the Ca^{2+} dependence of exocytosis observed in this study and indeed in mature mouse IHCs (Johnson et al., 2005, 2007). Therefore, another Ca^{2+} sensor could be involved and/or the expression levels of synaptic proteins such as SNAP25 and synaptobrevin 1 (Sendin et al., 2007) or the synaptic chaperone cysteine-string protein (Eybalin et al., 2002) could influence Ca^{2+} -exocytosis coupling. The expression of such molecules could be regulated by spontaneous AP activity present during early stages of development (Gu and Spitzer, 1995). Indeed, abnormal AP activity in prehearing IHCs prevented the developmental linearization in exocytotic Ca^{2+} dependence (Johnson et al., 2007). Therefore, in the maturing gerbil cochlea, the selective expression of synaptic molecules could be dictated by position-dependent differences in AP frequency. Although this has yet to be investigated, I_{Ca} in basal IHCs can support a higher AP frequency than apical cells (Johnson and Marcotti, 2008).

Kinetics of vesicle release in adult IHCs

At least two distinct kinetic components of vesicle release were evident in adult gerbil IHCs: a rapid component (RRP) and a slower but larger component (SRP). Although the SRP showed a nearly constant release rate up to at least 1.5 s in both cochlear regions, the RRP saturated with time constants of 52 ms (apical) and 42 ms (basal). Because I_{Ca} in IHCs exhibits very little inactivation for stimuli ≤ 100 ms (Johnson and Marcotti, 2008), the depletion of docked vesicles (apical, 534; basal, 619) was the most likely explanation for this initial component. The RRP in gerbil IHCs corresponded to ~ 26 (apical) and 28 (basal) vesicles per active zone, which was well correlated with anatomical observations (≤ 30 vesicles per synapse in mouse) (Khimich et al., 2005) and auditory afferent recordings (~ 12 vesicles per synapse in rat) (Goutman and Glowatzki, 2007).

IHC afferent synapses require rapid vesicle pool replenishment to continuously encode sound information. Although the release of vesicles from within the RRP was slightly reduced by repetitive stimulation, which was more evident in the apical region, IHCs could maintain exocytosis for prolonged time periods. Therefore, vesicle fusion and not their replenishment was the main rate limiting step for exocytosis, as also found in lower

vertebrates (Schnee et al., 2005). Rapid RRP replenishment has also been observed in guinea pig IHCs using two-photon imaging (Griesinger et al., 2005). In contrast, the SRP was more susceptible to pool depletion after repetitive stimulation, especially in low-frequency IHCs. Possible explanations for this could be a lower cytoplasmic vesicle density in apical cells and/or the poorer replenishment of ribbon-associated synaptic vesicles. Therefore, the synaptic machinery in adult mammalian IHCs appears to be specifically designed for sustaining neurotransmitter release with high-frequency cells being more indefatigable.

References

- Augustine GJ, Charlton MP (1986) Calcium dependence of presynaptic calcium current and postsynaptic response at the squid giant synapse. *J Physiol* 381:619–640.
- Augustine GJ, Charlton MP, Smith SJ (1985) Calcium entry and transmitter release at voltage-clamped nerve terminals of squid. *J Physiol* 367:163–181.
- Beurg M, Safieddine S, Roux I, Bouleau Y, Petit C, Dulon D (2008) Calcium- and otoferlin-dependent exocytosis by immature outer hair cells. *J Neurosci* 28:1798–1803.
- Beutner D, Moser T (2001) The presynaptic function of mouse cochlear inner hair cells during development of hearing. *J Neurosci* 21:4593–4599.
- Brandt A, Striessnig J, Moser T (2003) $\text{Ca}_v1.3$ channels are essential for development and presynaptic activity of cochlear inner hair cells. *J Neurosci* 23:10832–10840.
- Brandt A, Khimich D, Moser T (2005) Few $\text{Ca}_v1.3$ channels regulate the exocytosis of a synaptic vesicle at the hair cell ribbon synapse. *J Neurosci* 25:11577–11585.
- Brandt N, Kuhn S, Münkner S, Braig C, Winter H, Blin N, Vonthein R, Knipper M, Engel J (2007) Thyroid hormone deficiency affects postnatal spiking activity and expression of Ca^{2+} and K^+ channels in rodent inner hair cells. *J Neurosci* 27:3174–3186.
- Cody AR, Russell IJ (1987) The responses of hair cells in the basal turn of the guinea-pig cochlea to tones. *J Physiol* 383:551–569.
- Dallos P (1985) Response characteristics of mammalian cochlear hair cells. *J Neurosci* 5:1591–1608.
- Dodge FA Jr, Rahamimoff R (1967) Co-operative action of calcium ions in transmitter release at the muscular junction. *J Physiol* 193:419–432.
- Eybalin M, Renard N, Aure F, Safieddine S (2002) Cysteine-string protein in inner hair cells of the organ of Corti: synaptic expression and upregulation at the onset of hearing. *Eur J Neurosci* 15:1409–1420.
- Fettiplace R, Fuchs PA (1999) Mechanisms of hair cell tuning. *Annu Rev Physiol* 61:809–834.
- Fuchs PA (2005) Time and intensity coding at the hair cell's ribbon synapse. *J Physiol* 566:7–12.
- Glowatzki E, Fuchs PA (2002) Transmitter release at the hair cell ribbon synapse. *Nat Neurosci* 5:147–154.
- Goutman JD, Glowatzki E (2007) Time course and calcium dependence of transmitter release at a single ribbon synapse. *Proc Natl Acad Sci U S A* 104:16341–16346.
- Greenwood DD (1990) A cochlear frequency-position function for several species—29 years later. *J Acoust Soc Am* 87:2592–2605.
- Griesinger CB, Richards CD, Ashmore JF (2005) Fast vesicle replenishment allows indefatigable signalling at the first auditory synapse. *Nature* 435:212–215.
- Gu X, Spitzer NC (1995) Distinct aspects of neuronal differentiation encoded by frequency of spontaneous Ca^{2+} transients. *Nature* 375:784–787.
- Hackney CM, Mahendrasingam S, Penn A, Fettiplace R (2005) The concentration of calcium buffering proteins in mammalian cochlear hair cells. *J Neurosci* 25:7867–7875.
- Jahn R, Lang T, Südhof TC (2003) Membrane fusion. *Cell* 112:519–533.
- Jia S, Dallos P, He DZ (2007) Mechanoelectric transduction of adult inner hair cells. *J Neurosci* 27:1006–1014.
- Johnson SL, Marcotti W (2008) Biophysical properties of $\text{Ca}_v1.3$ calcium channels in gerbil inner hair cells. *J Physiol* 586:1029–1042.
- Johnson SL, Thomas MV, Kros CJ (2002) Membrane capacitance measurement using patch clamp with integrated self-balancing lock-in amplifier. *Pflugers Arch* 443:653–663.
- Johnson SL, Marcotti W, Kros CJ (2005) Increase in efficiency and reduc-

- tion in Ca^{2+} dependence of exocytosis during development of mouse inner hair cells. *J Physiol* 563:177–191.
- Johnson SL, Adelman JP, Marcotti W (2007) Genetic deletion of SK2 channels in mouse inner hair cells prevents the developmental linearization in the Ca^{2+} dependence of exocytosis. *J Physiol* 583:631–646.
- Keen EC, Hudspeth AJ (2006) Transfer characteristics of the hair cell's afferent synapse. *Proc Natl Acad Sci U S A* 103:5537–5542.
- Khimich D, Nouvian R, Pujol R, Tom Dieck S, Egner A, Gundelfinger ED, Moser T (2005) Hair cell synaptic ribbons are essential for synchronous auditory signalling. *Nature* 434:889–894.
- Kros CJ, Ruppertsberg JP, Rüsch A (1998) Expression of a potassium current in inner hair cells during development of hearing in mice. *Nature* 394:281–284.
- Lenzi D, Runyeon JW, Crum J, Ellisman MH, Roberts WM (1999) Synaptic vesicle populations in saccular hair cells reconstructed by electron tomography. *J Neurosci* 19:119–132.
- Linás R, Steinberg IZ, Walton K (1981) Relationship between presynaptic calcium current and postsynaptic potential in squid giant synapse. *Biophys J* 33:323–351.
- Marcotti W, Johnson SL, Holley MC, Kros CJ (2003a) Developmental changes in the expression of potassium currents of embryonic, neonatal and mature mouse inner hair cells. *J Physiol* 548:383–400.
- Marcotti W, Johnson SL, Rusch A, Kros CJ (2003b) Sodium and calcium currents shape action potentials in immature mouse inner hair cells. *J Physiol* 552:743–761.
- Marcotti W, Johnson SL, Kros CJ (2004) A transiently expressed SK current sustains and modulates action potential activity in immature mouse inner hair cells. *J Physiol* 560:691–708.
- Martinez-Dunst C, Michaels RL, Fuchs PA (1997) Release sites and calcium channels in hair cells of the chick's cochlea. *J Neurosci* 17:9133–9144.
- Moser T, Beutner D (2000) Kinetics of exocytosis and endocytosis at the cochlear inner hair cell afferent synapse of the mouse. *Proc Natl Acad Sci U S A* 97:883–888.
- Moser T, Neef A, Khimich D (2006) Mechanisms underlying the temporal precision of sound coding at the inner hair cell ribbon synapse. *J Physiol* 576:55–62.
- Müller M (1996) The cochlear place-frequency map of the adult and developing Mongolian gerbil. *Hear Res* 94:148–156.
- Mulroy MJ (1986) Patterns of afferent synaptic contacts in the alligator Lizard's cochlea. *J Comp Neurol* 248:263–271.
- Naraghi M, Neher E (1997) Linearized buffered Ca^{2+} diffusion in microdomains and its implications for calculation of $[\text{Ca}^{2+}]$ at the mouth of a calcium channel. *J Neurosci* 17:6961–6973.
- Neef A, Khimich D, Pirih P, Riedel D, Wolf F, Moser T (2007) Probing the mechanism of exocytosis at the hair cell ribbon synapse. *J Neurosci* 27:12933–12944.
- Neher E (1998) Vesicle pools and Ca^{2+} microdomains: new tools for understanding their roles in neurotransmitter release. *Neuron* 20:389–399.
- Nouvian R, Beutner D, Parsons TD, Moser T (2006) Structure and function of the hair cell ribbon synapse. *J Membr Biol* 209:153–165.
- Ohlemiller KK, Siegel JH (1994) Cochlear basal and apical differences reflected in the effects of cooling on responses of single auditory nerve fibers. *Hear Res* 80:174–190.
- Palmer AR, Russell IJ (1986) Phase-locking in the cochlear nerve of the guinea-pig and its relation to the receptor potential of inner hair-cells. *Hear Res* 24:1–15.
- Parsons TD, Lenzi D, Almers W, Roberts WM (1994) Calcium-triggered exocytosis and endocytosis in an isolated presynaptic cell: capacitance measurements in saccular hair cells. *Neuron* 13:875–883.
- Platzer J, Engel J, Schrott-Fischer A, Stephan K, Bova S, Chen H, Zheng H, Striessnig J (2000) Congenital deafness and sinoatrial node dysfunction in mice lacking class D L-type Ca^{2+} channels. *Cell* 102:89–97.
- Roberts WM, Jacobs RA, Hudspeth AJ (1990) Colocalization of ion channels involved in frequency selectivity and synaptic transmission at presynaptic active zones of hair cells. *J Neurosci* 10:3664–3684.
- Roux I, Safieddine S, Nouvian R, Grati M, Simmler MC, Bahloul A, Perfettini I, Le Gall M, Rostaing P, Hamard G, Triller A, Avan P, Moser T, Petit C (2006) Otoferlin, defective in a human deafness form, is essential for exocytosis at the auditory ribbon synapse. *Cell* 127:277–289.
- Ryan A (1976) Hearing sensitivity of the Mongolian gerbil, *Meriones unguiculatus*. *J Acoust Soc Am* 5:1222–1226.
- Schmitz F, Königstorfer A, Südhof TC (2000) RIBEYE, a component of synaptic ribbons: a protein's journey through evolution provides insight into synaptic ribbon function. *Neuron* 28:857–872.
- Schnee ME, Lawton DM, Furness DN, Benke TA, Ricci AJ (2005) Auditory hair cell-afferent fiber synapses are specialized to operate at their best frequencies. *Neuron* 47:243–254.
- Schug N, Braig C, Zimmermann U, Engel J, Winter H, Ruth P, Blin N, Pfister M, Kalbacher H, Knipper M (2006) Differential expression of otoferlin in brain, vestibular system, immature and mature cochlea of the rat. *Eur J Neurosci* 24:3372–3380.
- Sendin G, Bulankina AV, Riedel D, Moser T (2007) Maturation of ribbon synapses in hair cells is driven by thyroid hormone. *J Neurosci* 27:3163–3173.
- Slepecky NB, Galsky MD, Swartzentruber-Martin H, Savage J (2000) Study of afferent nerve terminals and fibers in the gerbil cochlea: distribution by size. *Hear Res* 144:124–134.
- Sobkowicz HM, Rose JE, Scott GE, Slapnick SM (1982) Ribbon synapses in the developing intact and cultured organ of Corti in the mouse. *J Neurosci* 2:942–957.
- Sterling P, Matthews G (2005) Structure and function of ribbon synapses. *Trends Neurosci* 28:20–29.
- Thoreson WB, Rabl K, Townes-Anderson E, Heidelberger R (2004) A highly Ca^{2+} -sensitive pool of vesicles contributes to linearity at the rod photoreceptor ribbon synapse. *Neuron* 42:595–605.
- Tucker T, Fettiplace R (1995) Confocal imaging of calcium microdomains and calcium extrusion in turtle hair cells. *Neuron* 15:1323–1335.
- von Gersdorff H, Vardi E, Matthews G, Sterling P (1996) Evidence that vesicles on the synaptic ribbon of retinal bipolar neurons can be rapidly released. *Neuron* 16:1221–1227.
- Woolf NK, Ryan AF (1984) The development of auditory function in the cochlea of the mongolian gerbil. *Hear Res* 13:277–283.
- Yasunaga S, Grati M, Cohen-Salmon M, El-Amraoui A, Mustapha M, Salem N, El-Zir E, Loiselet J, Petit C (1999) A mutation in OTOF, encoding otoferlin, a FER-1-like protein, causes DFNB9, a nonsyndromic form of deafness. *Nat Genet* 21:363–369.