

Calcium- and Otoferlin-Dependent Exocytosis by Immature Outer Hair Cells

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Immature cochlear outer hair cells (OHCs) make transient synaptic contacts (ribbon synapses) with type I afferent nerve fibers, but direct evidence of synaptic vesicle exocytosis is still missing. We thus investigated calcium-dependent exocytosis in murine OHCs at postnatal day 2 (P2)–P3, a developmental stage when calcium current maximum amplitude was the highest. By using time-resolved patch-clamp capacitance measurements, we show that voltage step activation of L-type calcium channels triggers fast membrane capacitance increase. Capacitance increase displayed two kinetic components, which are likely to reflect two functionally distinct pools of synaptic vesicles, a readily releasable pool (RRP; $\tau = 79$ ms) and a slowly releasable pool ($\tau = 870$ ms). The RRP size and maximal release rate were estimated at ~ 1200 vesicles and $\sim 15,000$ vesicles/s, respectively. In addition, we found a linear relationship between capacitance increase and calcium influx, like in mature inner hair cells (IHCs). These results give strong support to the existence of efficient calcium-dependent neurotransmitter release in immature OHCs. Moreover, we show that immature OHCs, just like immature IHCs, are able to produce regenerative calcium-dependent action potentials that could trigger synaptic exocytosis *in vivo*. Finally, the evoked membrane capacitance increases were abolished in P2–P3 OHCs from mutant *Otof*^{-/-} mice defective for otoferlin, despite normal calcium currents. We conclude that otoferlin, the putative major calcium sensor at IHC ribbon synapses, is essential to synaptic exocytosis in immature OHCs too.

Key words: cochlea; organ of Corti; outer hair cells; ribbon synapses; exocytosis; otoferlin

Introduction

The sense of hearing depends on fast, graded neurotransmitter release at the ribbon synapses of cochlear inner hair cells (IHCs) (for review, see Fuchs, 2005). The finely tuned and high-rate neurotransmitter release at IHC synapses involves synaptic vesicle fusion with the plasma membrane, which is precisely controlled by calcium influx through L-type (Ca_v1.3) calcium channels (Brandt et al., 2003, 2005; Johnson et al., 2005). The molecular events that trigger synaptic exocytosis in IHCs are still poorly understood. These cells apparently lack some major proteins found at CNS synapses, including the calcium sensor synaptotagmin I (Safieddine and Wenthold, 1999). Otoferlin, a large synaptic vesicle transmembrane protein with six C2 domains, has been proposed to be the major calcium sensor at the IHC ribbon synapse (Roux et al., 2006). Otoferlin indeed interacts with the SNARE (soluble N-ethylmaleimide-sensitive factor attachment protein receptor) proteins, syntaxin-1 and synaptosome-

associated protein of 25 kDa (SNAP-25), in a calcium-dependent way. Furthermore, in deaf mice carrying null otoferlin alleles (*Otof*^{-/-} mice), IHCs lack calcium-evoked-exocytosis (Roux et al., 2006).

In contrast to IHCs, outer hair cells (OHCs) do not significantly contribute to the afferent signaling, but rather act as sound signal amplifiers in the mature cochlea (for review, see Dallos et al., 2006). In immature OHCs, however, otoferlin is present during the first postnatal days in the mouse (Roux et al., 2006; Schug et al., 2006). At this stage, OHCs are contacted by type-I afferent nerve fibers (Pujol, 1985; Echterler, 1992; Huang et al., 2007) that bear glutamate receptors (Knipper et al., 1997; Engel et al., 2006), and they have presynaptic ribbon structures (Sobkowicz et al., 1982), which suggests that transient synaptic exocytosis takes place in these cells. In addition, immature OHCs express Ca_v1.3 calcium currents (Michna et al., 2003) that resemble those controlling synaptic exocytosis in mature IHCs. Notably, Knirsch et al. (2007) have found that Ca_v1.3 channels persist in a subpopulation of mature OHCs located at the cochlear apical end. These cells still have presynaptic ribbons and afferent type-I fibers (Pujol et al., 1997), and they express syntaxin-1 and SNAP-25 (Safieddine and Wenthold, 1999) as well as otoferlin (Roux et al., 2006), suggesting that they too undergo calcium-dependent synaptic exocytosis. The existence of calcium-dependent synaptic exocytosis, however, has so far not been demonstrated in mature or immature OHCs.

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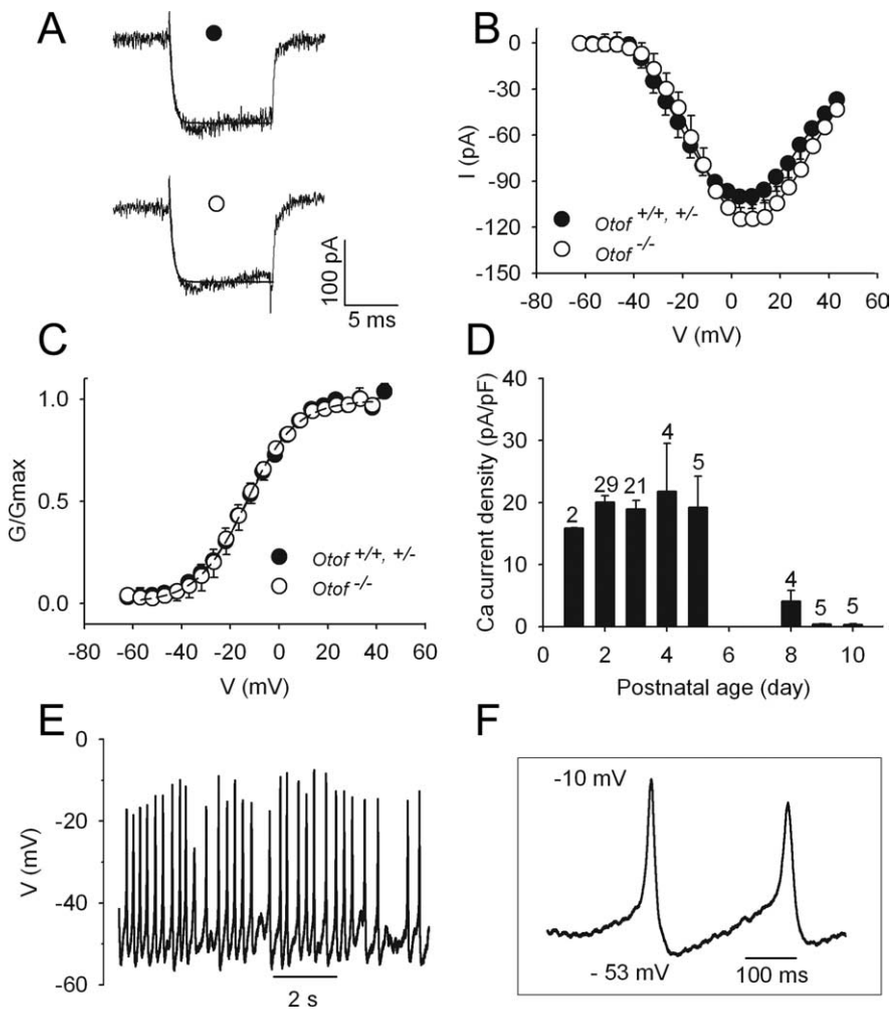


Figure 1. Properties of calcium currents in immature OHCs from *Otof*^{+/+/+} and *Otof*^{-/-} mice. **A**, Examples of I_{Ca} recorded in OHCs of *Otof*^{+/+/+} (●) and *Otof*^{-/-} (○) P2 mice. Currents were evoked by 10 ms voltage steps from a -82 mV holding potential to 0 mV. Single-exponential fits (solid lines) show similar activation kinetics, with $\tau = 342 \mu\text{s}$ (●) and $340 \mu\text{s}$ (○). **B**, Comparative I_{Ca} - V curves in OHCs of *Otof*^{+/+/+} and *Otof*^{+/-} (●; $n = 11$) and *Otof*^{-/-} (○; $n = 9$) mice. **C**, Average normalized I_{Ca} conductance in *Otof*^{+/+/+}, *Otof*^{+/-}, and *Otof*^{-/-} mice. Continuous lines are Boltzmann fits with the following parameters: $G_{\text{max}} = 2.6 \pm 0.4$ and 3 ± 0.2 nS; $V_{1/2} = -12.2 \pm 0.5$ and -12.3 ± 0.5 mV; and $k = 10.1 \pm 0.4$ and 9.9 ± 0.5 mV for *Otof*^{+/+/+} and *Otof*^{+/-} and *Otof*^{-/-} OHCs, respectively. **D**, Peak calcium current density (expressed in pA/pF) as a function of postnatal age in *Otof*^{+/+/+} and *Otof*^{+/-} OHCs. The numbers of tested cells are indicated on the histogram. Note the drastic decrease of I_{Ca} density after P6. **E**, Example of a train of spontaneous action potentials recorded from a P3 *Otof*^{+/+/+} OHC. The voltage recording was made in the absence of any current injection, in artificial perilymph with 10 mM calcium, at RT. **F**, Enlarged view of two action potentials.

Using cell membrane capacitance measurements, we report that calcium-dependent exocytosis actually occurs in immature OHCs. Moreover, we found that this exocytosis was absent in OHCs from otoferlin-deficient mice. Finally, we were able to record spontaneous, calcium-dependent action potentials in immature OHCs, which may account for synaptic vesicle release *in vivo*.

Materials and Methods

Animals. Experiments were performed on mice obtained by interbreeding of *Otof*^{+/-} mice (Roux et al., 2006). The overall study was performed on 66 mice (15 *Otof*^{+/+/+}, 32 *Otof*^{+/-}, and 19 *Otof*^{-/-}) issued from 36 different litters. All experiments were performed in accordance with the European Community Council Directive 86/609/EEC.

Immunohistochemistry. Tissue processing, immunohistochemistry, and confocal imaging were performed as described previously (Roux et al., 2006). Whole-mount preparations of postnatal day 3 (P3)

organ of Corti were double stained for otoferlin and ribeye using a monoclonal antibody (1/500) and a polyclonal antibody (1/500), respectively.

Whole-cell recordings. Experiments were performed on mouse IHCs and OHCs from isolated organ of Corti apical coils, ranging from P1 to P10. Extracellular solution contained the following (in mM): 115 NaCl, 6 KCl, 10 CaCl₂, 30 TEA, 2 Na pyruvate, 8 glucose, and 10 Na-HEPES, pH 7.4. Tetrodotoxin ($1 \mu\text{M}$), apamin ($1 \mu\text{M}$), and XE-991 ($1 \mu\text{M}$) (Sigma-Aldrich, St. Quentin Fallavier, France) were added to the extracellular solution to block Na⁺ and K⁺ conductances. Because we worked at room temperature (RT; 22–24°C), we used, unless otherwise specified, 10 mM CaCl₂ in the extracellular solution to increase the calcium current (I_{Ca}). Recording pipette solution contained the following (in mM): 130 Cs-gluconate, 1 MgCl₂, 5 Na₂ATP, 0.5 Na₂GTP, 5 TEA, 1 EGTA, and 10 Cs-HEPES, pH 7.2. Spontaneous action potentials (APs), however, were recorded using a potassium-based intracellular solution (in mM: 135 KCl, 0.1 CaCl₂, 1.5 MgCl₂, 11 EGTA, 5 HEPES, and 2.5 Na₂ATP, pH 7.4) and the above-described extracellular solution with no pharmacological blocker.

Capacitance measurement. Changes in cell membrane capacitance (ΔC_m) were used to monitor fusion of vesicles with the plasma membrane during exocytosis, and measured using the tracking circuitry of an Optopatch amplifier as described by Johnson et al. (2002) (for details, see supplemental material, available at www.jneurosci.org).

Data analysis. Curve fitting and analysis were done using Origin software (OriginLab, Northampton, MA) (see supplemental material, available at www.jneurosci.org). The criterion for statistical significance was chosen to be $p < 0.05$ and evaluated by Student's t tests. Variability is reported as \pm SEM.

Results

Calcium currents and spontaneous action potentials in immature OHCs

Neonatal OHCs displayed rapidly activating I_{Ca} in response to depolarizing voltage steps (Fig. 1A). Analysis of the calcium current–voltage relationship indicated that I_{Ca} activates near -40 mV and reaches a maximum at $+5$ mV (Fig. 1B). I_{Ca} was not significantly different between *Otof*^{+/+/+} and *Otof*^{+/-} ($I_{\text{max}} = 100.1 \pm 7.0$ pA; $n = 11$) and *Otof*^{-/-} ($I_{\text{max}} = 114.8 \pm 7.5$ pA; $n = 9$) OHCs. Normalized conductance–voltage curves also gave indistinguishable parameters (G_{max} , $V_{1/2}$, and k) between *Otof*^{+/+/+} and *Otof*^{+/-} ($G_{\text{max}} = 2.6 \pm 0.4$ nS; $V_{1/2} = -12.2 \pm 0.5$ mV; $k = 10.1 \pm 0.4$ mV) and *Otof*^{-/-} ($G_{\text{max}} = 3.0 \pm 0.2$ nS; $V_{1/2} = -12.3 \pm 0.5$ mV; $k = 9.9 \pm 0.5$ mV) cells when fitted with a first-order Boltzmann equation (Fig. 1C). The mean cell capacitance was not significantly different between P2–P3 *Otof*^{+/+/+} and *Otof*^{+/-} OHCs (6.0 ± 0.2 pF; $n = 50$) and P2–P3 *Otof*^{-/-} OHCs (5.9 ± 0.2 pF; $n = 20$) either. In *Otof*^{+/+/+} and *Otof*^{+/-} OHCs, the calcium current density was maximal between P2 and P5, and drastically decreased after P8 (Fig. 1D). This result is in good agreement with previous studies by Michna et al. (2003) and Knirsch et al. (2007). A similar large decrease of I_{Ca}

was observed in *Otof*^{-/-} OHCs (Fig. 1D). This result is in good agreement with previous studies by Michna et al. (2003) and Knirsch et al. (2007). A similar large decrease of I_{Ca}

occurred in *Otof*^{-/-} OHCs between P2–P3 ($n = 29$) and P10 ($n = 4$) (data not shown).

Regenerative APs could be recorded in all P3 OHCs studied, in 10 mM extracellular calcium (five *Otof*^{+/+} and *Otof*^{+/-} and three *Otof*^{-/-} OHCs). Spontaneous AP firing was observed without injecting current, from a mean resting membrane potential of -50 ± 1 mV (range, -46 to -56 mV; $n = 8$). Spontaneous APs occurred at a mean frequency of 1.7 ± 0.6 Hz, and had a maximum peak amplitude of 33 ± 4 mV and a mean half-width of 29 ± 6 ms (Fig. 1E, F). Similar AP firing was also recorded in five additional P3 (*Otof*^{+/-}) OHCs in artificial perilymph containing 1.3 mM CaCl₂ at near-physiological temperature ($34 \pm 2^\circ\text{C}$). These APs are likely to be driven by calcium currents, because they were insensitive to tetrodotoxin, and were blocked by 50 μM of the L-type calcium channel antagonist nifedipine. Moreover, APs were not observed in P9 OHCs ($n = 4$; data not shown), when only very low I_{Ca} could be recorded (Fig. 1D). These results are consistent with a previous report by Marcotti and Kros (1999), showing voltage responses to current injection that suggested regenerative APs in neonatal mouse OHCs.

Calcium-dependent synaptic exocytosis

To demonstrate calcium-dependent exocytosis in OHCs, we recorded ΔC_m during voltage-step activation of I_{Ca} . We used P2–P3 immature OHCs to avoid the prestin electromotility contribution to cell membrane capacitance variation (Abe et al., 2007). Indeed, fast and reversible non-linear capacitance changes caused by the voltage activation of prestin were absent in apical OHCs before P4 (data not shown).

At P2–P3, OHCs display large I_{Ca} (Fig. 1D). Calcium entry induced by a 100 ms voltage step from -80 to 0 mV produced a mean ΔC_m of 34.5 ± 7.3 fF ($n = 33$, 8 *Otof*^{+/+} and 25 *Otof*^{+/-} OHCs). An example of C_m recording is shown in Fig. 2A. The C_m jump was immediate, and the maximum value was already reached at the end of the 100 ms depolarizing step. The C_m increase was sustained over a period of several hundred milliseconds after the end of the stimulation. Analysis of the ΔC_m –voltage relationship showed a positive correlation between ΔC_m and I_{Ca} amplitude (Fig. 2B). To verify that exocytosis was indeed triggered by a rise in intracellular calcium concentration, some OHCs were recorded using an intrapipette solution containing 10 mM of the fast Ca²⁺-chelator BAPTA (instead of 1 mM EGTA). In the 10 mM BAPTA condition, OHCs, while showing a large peak I_{Ca} of 67 ± 5.3 pA, only showed very small C_m responses, averaging 2.2 ± 0.8 fF ($n = 6$, 2 *Otof*^{+/+} and 4 *Otof*^{+/-}), to cell depolarization. In addition, ΔC_m responses, recorded in the EGTA condition, were completely abolished when I_{Ca} was blocked by nifedipine (50 μM) ($n = 5$ cells; data not shown). Together, these results give evidence of synaptic exocytosis in immature OHCs and the involvement of L-type calcium channels

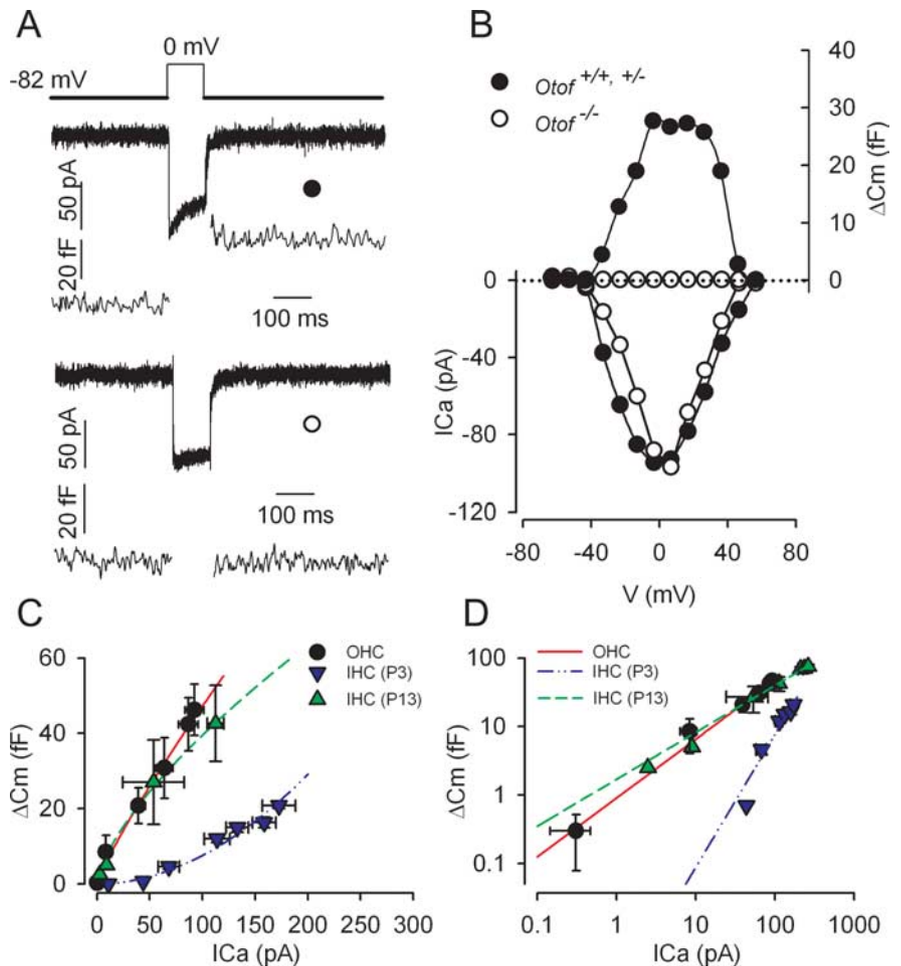


Figure 2. Calcium-evoked exocytosis in OHCs and IHCs from *Otof*^{+/+} and *Otof*^{-/-} mice. **A**, Examples of I_{Ca} with the corresponding ΔC_m response in *Otof*^{+/+} (●) and *Otof*^{-/-} (○) OHCs at P2. Cells were stimulated by a 100 ms voltage step from a holding potential of -82 to 0 mV. **B**, Voltage– I_{Ca} relationship (bottom) and corresponding ΔC_m (top) are shown for the same cells as in **A**. **C, D**, Comparative I_{Ca} dependence of ΔC_m in P2–P3 OHCs (●; $n = 5$), P3 IHCs (▼; $n = 6$), and P13 IHCs (▲; $n = 3$) from *Otof*^{+/+} and *Otof*^{-/-} mice. Increasing I_{Ca} was elicited by stepping the cells from a -82 mV holding potential to different potentials up to 0 mV, in 10 mV increments (100 ms step duration). ΔC_m was plotted against corresponding I_{Ca} . The synaptic transfer function was best fitted with a power function $Y = aX^N$, with $a = 1, 0.002$, and 1.3 and $N = 0.84, 1.9$, and 0.68 in P2–P3 OHCs, P3 IHCs, and P13 IHCs, respectively. **D**, I_{Ca} dependence of ΔC_m as in **C** but plotted in double-logarithmic scales.

in this process. Increases in C_m and I_{Ca} could also be recorded in more physiological conditions at $34 \pm 2^\circ\text{C}$ with 1.3 mM external CaCl₂ (data not shown). A calcium efficiency (ratio $\Delta C_m/I_{\text{Ca}}$ –(peak)) of 0.35 fF/pA ($n = 3$) was recorded in these conditions. This value is similar to the efficiency we recorded in the 10 mM CaCl₂ condition, at RT (0.31 fF/pA; $n = 33$).

To further characterize the calcium-dependent exocytosis in immature OHCs, we studied the relationship between calcium inflow and exocytosis, i.e., the synaptic transfer function. P2–P3 OHCs were stimulated by a 100 ms voltage step to a range of potentials varying between -50 and 0 mV, and ΔC_m responses were plotted against corresponding I_{Ca} amplitudes. Peak I_{Ca} was preferred to I_{Ca} integral (total amount of calcium charges) because of the frequent presence of a residual outward potassium current carried by SK channels incompletely blocked by apamin. The relationship between ΔC_m and I_{Ca} was best fitted by using a power function, with $N = 0.84$ (Fig. 2C). For comparison, the synaptic transfer function was also studied in P3 IHCs and “nearly mature” (P13) IHCs, in similar 10 mM CaCl₂ conditions (Fig. 2C, D). Interestingly, P2–P3 OHC exocytosis had a calcium efficiency similar to that of P13 IHCs (0.30 fF/pA and power

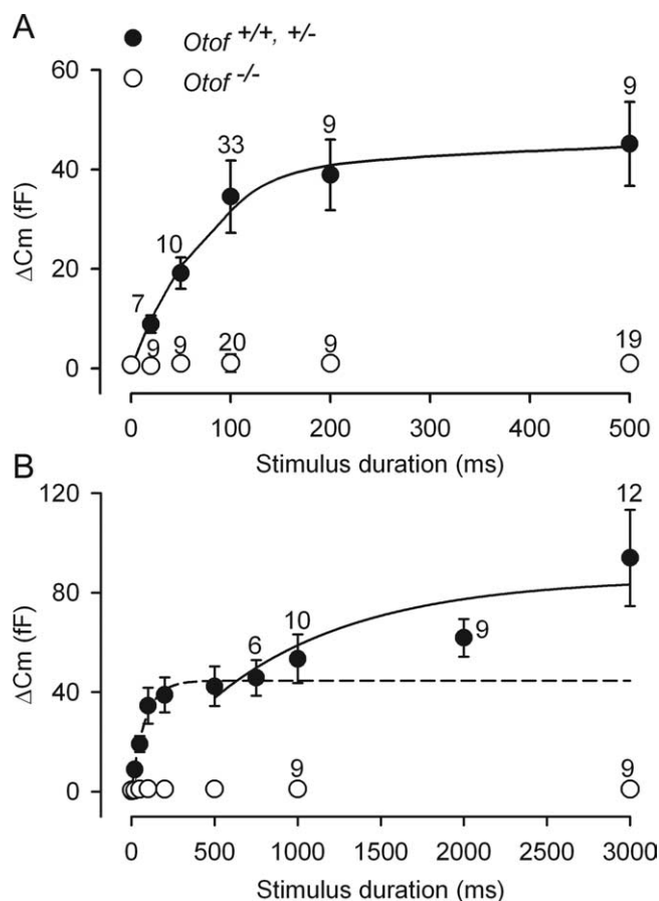


Figure 3. Kinetics of exocytosis in immature OHCs. *A, B*, Average ΔC_m was plotted as a function of the voltage stimulus duration. All cells were voltage stepped to 0 mV from a holding potential of -82 mV. ΔC_m was obtained in response to stimuli with varying durations, up to 500 ms (*A*) or 3000 ms (*B*). Solid lines correspond to two consecutive single-exponential fits, with $\tau = 79.3$ ms in *A*, and $\tau = 870$ ms in *B*, which define an RRP and an SRP of synaptic vesicles, respectively. In *B*, the dashed line is extrapolated from the fit shown in *A*. Numbers of tested P2–P3 OHCs are indicated on the graphs.

function fitted with $N = 0.7$), whereas that of P3 IHCs was lower (0.11 fF/pA and power function fitted with $N \sim 2$).

Kinetics of synaptic vesicle release

We used voltage steps to 0 mV with varying durations to study the kinetics of vesicle release in P2–P3 OHCs. This method has been used to show the existence of two different pools of vesicles in IHC synaptic exocytosis (Moser and Beutner, 2000; Beutner et al., 2001; Johnson et al., 2005). In these studies, two components of vesicular release have been observed. The fast ΔC_m component, which can be recorded during brief stimuli (< 500 ms), is believed to account for exocytosis of the ready releasable pool of vesicles (RRP), whereas the slower ΔC_m component, which can be measured during longer depolarizations, would be caused by the release of a second pool of vesicles [slowly releasable pool (SRP)], located at a larger distance from the calcium entry site.

A fast exocytosis component was characterized in P2–P3 OHCs (Fig. 3*A*). Whereas short stimuli (5–10 ms) were unable to elicit ΔC_m responses above the background noise (1.8 ± 1.0 fF), the amplitude of ΔC_m responses rapidly increased with stimulus duration between 20 and 100 ms, and reached a steady state value at 500 ms. A single-exponential fit with a time constant of 79.3 ms and a maximal ΔC_m of 44.5 fF sug-

gested that this fast exocytosis component involves a single RRP. Assuming a ΔC_m of 37 aF per vesicle (Lenzi et al., 1999), calculation gave a RRP of ~ 1200 vesicles in OHCs. This RRP could be released with a maximum rate of $\sim 15,100$ vesicles/s (560 fF/s). This value is similar to the release rate reported by Johnson et al. (2005) in P10–P20 mouse IHCs maintained at body temperature in physiological (1.3 mM) extracellular calcium (rate constant of 459 fF/s with $\tau = 53$ ms). The RRP of immature OHCs, however, is significantly larger than the RRP reported in mature IHCs, i.e., 660 synaptic vesicles or less (Johnson et al., 2005). Intriguingly enough, hair cells from the frog sacculle and the turtle auditory papilla have both larger RRP responses and faster kinetics of exocytosis (Edmonds et al., 2004; Schnee et al., 2005). This may suggest different mechanisms for synaptic vesicle fusion in lower vertebrates.

During longer stimulations, a second, slower exocytosis component (SRP) could be identified in neonatal P2–P3 OHCs. This component was also best fitted using a single exponential starting after 500 ms ($\tau = 870$ ms; maximal $\Delta C_m = 86$ fF), giving a maximal release rate of ~ 2670 vesicles/s (99 fF/s) (Fig. 3*B*). Notably, the C_m responses over the entire time range (up to 3 s) could not be accurately fitted with a double exponential. A good fit for the RRP and SRP was obtained using two sequential and independent single-exponential functions, which argues for a delayed activation of the second phase of exocytosis. The physiological significance of this secondary release, however, remains questionable because it was observed during nonphysiological long depolarization. Whether the SRP release also occurs at the ribbon synapses or involves other nonsynaptic release sites remains to be determined.

Otoferlin is required for calcium-dependent exocytosis in immature OHCs

At P3, otoferlin is abundant both in OHCs and IHCs (Roux et al., 2006). Otoferlin was detected throughout the cytoplasm, but the immunostaining was stronger in the basolateral membrane area, where the ribbon synapses are located (Fig. 4). Indeed, the otoferlin distribution largely overlapped with that of the integral ribbon protein ribeye (Schmitz et al., 2000) in both cell types (Fig. 4, top). We thus asked whether synaptic exocytosis in immature OHCs depends on the presence of otoferlin, by studying the ΔC_m response to cell depolarization in P2–P3 *Otof*^{-/-} mice that lack otoferlin. In *Otof*^{-/-} OHCs, we found that the exocytosis response was abolished (ΔC_m responses below background threshold of 1.8 fF; $n = 20$) (Fig. 2*A, B*), despite I_{Ca} values similar to wild-type mice (maximum current density of 17.1 ± 1.0 pA/pF in *Otof*^{-/-} mice compared with 18.5 ± 1.3 pA/pF in *Otof*^{+/-} and *Otof*^{+/+} mice) (Fig. 1*A–C*) and apparent normal presynaptic active zone assembly, as suggested by the ribeye immunostaining (Fig. 4, bottom). Longer depolarizations (up to several seconds) that allow larger intracellular calcium increase (Beutner et al., 2001) were ineffective as well (Fig. 3*B*). We conclude that otoferlin is required for both RRP and SRP synaptic vesicle exocytosis in immature OHCs.

Discussion

Our results provide strong evidence for the existence of spontaneous calcium action potentials and synaptic exocytosis in P2–P3 mouse OHCs that recording of EPSCs will definitively establish. Interestingly, P2–P3 OHC exocytosis had a calcium efficiency similar to that of mature P13 IHCs, whereas that of

immature P3 IHCs was three times lower. This is in good agreement with morphological studies at early postnatal stages, showing that OHCs contain more ribbons than IHCs (Sobkowicz et al., 1982). Johnson et al. (2005), using a physiological (1.3 mM) extracellular calcium condition, also found a lower efficiency of exocytosis in the developing IHCs maintained at body temperature. The almost linear, noncooperative relationship between ΔC_m and I_{Ca} in mature P13 IHCs and P2–P3 OHCs suggests that exocytosis relies on a one-to-one relationship between calcium binding and vesicle fusion events. A similar calcium dependence of release has been found in hair cells from the turtle auditory papilla (power function fitted with $N = 0.9$) (Schnee et al., 2005). Furthermore, by recording postsynaptic currents, Keen and Hudspeth (2006) and Goutman and Glowatzki (2007) also found a linear calcium dependence of transmitter release at the level of a single hair cell ribbon synapse. Interestingly, our results demonstrate that otoferlin, the putative calcium sensor at IHC ribbon synapses (Roux et al., 2006), is essential to synaptic exocytosis in immature OHCs too.

We propose that synaptic activity in the immature OHCs contributes to the spontaneous discharge activity recorded in spiral ganglion neurons before the onset of hearing (Jones et al., 2007). Although electrical spiking activity is not essential to the synapse formation (Verhage et al., 2000), it is well established as critically involved in the maturation of neuronal circuitry (for review, see Hua and Smith, 2004). Synchronized spontaneous rhythmic activity of neurotransmission has indeed an instructive role in the refinement of neuronal connections in the developing retina (Galli and Maffei, 1988; Torborg and Feller, 2005). We suggest that the spontaneous spiking activity of immature OHC ribbon synapses plays a similar role in the developing auditory system.

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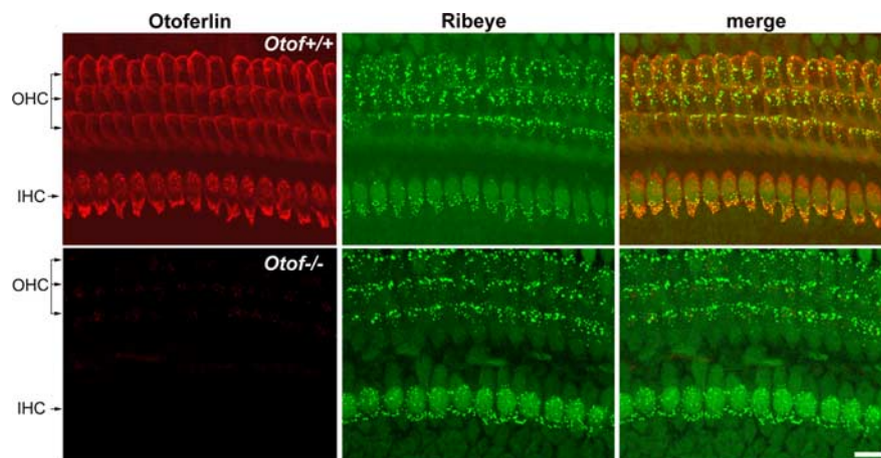


Figure 4. Otoferlin and ribeye distributions in the organ of Corti of P3 *Otof*^{+/+} and *Otof*^{-/-} mice. In the P3 *Otof*^{+/+} mouse, otoferlin (red) is detected both in IHCs and OHCs. At this stage, both cell types contain synaptic ribbons, revealed by the ribeye immunolabeling (green). The otoferlin and ribeye stainings largely overlap, as shown in yellow in the merged panel. In the P3 *Otof*^{-/-} mouse, otoferlin is not detected in hair cells, whereas the ribeye immunolabeling is undistinguishable from that of *Otof*^{+/+} mice. Scale bar, 10 μ m.

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