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Control of Exocytosis by Synaptotagmins and Otoferlin in Auditory Hair Cells

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In pre-hearing mice, vesicle exocytosis at cochlear inner hair cell (IHC) ribbon synapses is triggered by spontaneous Ca\(^{2+}\) spikes. At the onset of hearing, IHC exocytosis is then exclusively driven by graded potentials, and is characterized by higher Ca\(^{2+}\)-efficiency and improved synchronization of vesicular release. The molecular players involved in this transition are still unknown. Here we addressed the involvement of synaptotagmins and otoferlin as putative Ca\(^{2+}\) sensors in IHC exocytosis during postnatal maturation of the cochlea.

Using cell capacitance measurements, we showed that Ca\(^{2+}\)-evoked exocytosis in mouse IHCs switches from an otoferlin-independent to an otoferlin-dependent mechanism at postnatal day 4. During this early exocytotic period, several synaptotagmins (Syts), including Syt1, Syt2 and Syt7, were detected in IHCs. The exocytotic response as well as the release of the readily releasable vesicle pool (RRP) was, however, unchanged in newborn mutant mice lacking Syt1, Syt2 or Syt7 (Syt1\(^{-/-}\), Syt2\(^{-/-}\), and Syt7\(^{-/-}\) mice). We only found a defect in RRP recovery in Syt1\(^{-/-}\) mice which was apparent as a strongly reduced response to repetitive stimulations. In post-hearing Syt2\(^{-/-}\) and Syt7\(^{-/-}\) mutant mice, IHC synaptic exocytosis was unaffected. The transient expression of Syt1 and Syt2, which were no longer detected in IHCs after the onset of hearing, indicates that these two most common Ca\(^{2+}\)-sensors in CNS synapses are not involved in mature IHCs. We suggest that otoferlin underlies highly efficient Ca\(^{2+}\)-dependent membrane-membrane fusion, a process likely essential to increase the probability and synchrony of vesicle fusion events at the mature IHC ribbon synapse.

Introduction

Before the onset of hearing, cochlear inner hair cells (IHCs) exhibit spontaneous Ca\(^{2+}\) spiking activity that triggers synaptic exocytosis (Kros et al., 1998; Beutner et al., 2001; Marocci et al., 2003; Tritsch and Bergles, 2010) and postsynaptic trains of action potentials (APs) in cochlear ganglion neurons (Lippe, 1994; Jones et al., 2007; Tritsch et al., 2007). In mature IHCs, exocytosis is then driven by sound-evoked fast graded receptor potentials. In both immature and mature IHCs, neurotransmitter release is triggered by Ca\(^{2+}\) influx flowing through L-type Ca\(^{2+}\) channels (Moser and Beutner, 2000; Glowatzki and Fuchs, 2002; Brandt et al., 2003), and the efficiency of brief Ca\(^{2+}\) currents in evoking release increases with cell maturation (Beutner and Moser, 2001; Johnson et al., 2005). Synchronization of the multivesicular release process (Glowatzki and Fuchs, 2002) also improves with IHC maturation (Grant et al., 2010), which suggests that concomitant, as yet poorly understood changes take place in the IHC synaptic machinery.

Like in most neurosecretory cells, synaptic exocytosis in IHCs is thought to involve interactions of SNARE (soluble N-ethylmaleimide-sensitive-factor attachment protein receptor) complex proteins attached to synaptic vesicles (synaptobrevin) with the target plasma membrane (SNAP-25 and syntaxin 1) (Safieddine and Wenthold, 1999). Synaptotagmins (Syts), a large family of transmembrane proteins containing tandem Ca\(^{2+}\)-binding C2-domains, confer Ca\(^{2+}\) sensitivity to SNARE-dependent vesicle fusion in the CNS (Chapman, 2008), but their implication in IHC synaptic exocytosis is still unclear. Indeed, the transcripts encoding Syt1 and Syt2, the major Ca\(^{2+}\) sensors for fast synchronized transmitter release in central neurons (Geppert et al., 1994; Sun et al., 2007), have not been detected in mature IHCs (Safieddine and Wenthold, 1999). Therefore, otoferlin, a six C2-domain transmembrane protein (Yasunaga et al., 1999), has been proposed as a major Ca\(^{2+}\) sensor required for Ca\(^{2+}\)-evoked exocytosis in IHCs (Roux et al., 2006).
The synaptic active zone of IHCs has been proposed to be organized in independent Ca2+ nanodomains, where the activity of one or few Ca2+ channels is sufficient to activate release of a nearby vesicle (Brandt et al., 2005), allowing the synapse to function in a linear regime (Johnson et al., 2005; Keen and Hudspeth, 2006; Goutman and Glowatzki, 2007). Recently, Syt4, which does not bind Ca2+ and is not established as a Ca2+ sensor (von Poser et al., 1997), has been shown to be required for the linear Ca2+ dependence of exocytosis (Johnson et al., 2010). Notably, this latter study also reported the expression of Syt1 and Syt2 in immature and mature IHCs, respectively, but the functional implication of these findings has not yet been investigated. To clarify the respective roles of these putative Ca2+ sensors at the IHC ribbon synapse along the course of cochlear maturation, we studied their expression and analyzed Ca2+ dependent exocytosis in IHCs from Otof−/−, Syt1−/−, Syt2−/− and Syt7−/− mutant mice that lack otoferlin, Syt1, Syt2 and Syt7, respectively (Geppert et al., 1994; Roux et al., 2006; Sun et al., 2007).

Materials and Methods

Electrophysiological recordings

Experiments were performed on mice, both male and female, obtained by interbreeding of heterozygous mutant mice (Otof−/− or Syt−/− animals) and genotyped using PCR. Recordings and analyses were performed in blind before knowing the genotype of the mice. All experiments were performed in accordance with the European Communities Council Directive of November 24, 1986 (86/609/EEC). All cases were taken to minimize animals’ pain. Whole-cell recordings were performed on excised apical coils of organs of Corti as previously described (Beurg et al., 2008). Dishes were done in a solution containing the following (in mM): 135 NaCl, 5.8 KCl, 1.3 CaCl2, 0.9 MgCl2, 0.9 NaH2PO4, 5.6 glucose, 2 Na-pyruvate, 10 HEPES, pH 7.4.

Borosilicate patch electrodes (World Precision Instruments) were coated with ski wax (SWIX) to minimize electrode capacitance. Pipette resistances in solution were 2–3 MΩ. Experiments were done at room temperature (20–22°C).

Recordings in otoferlin-null mice were acquired with an Optopatch amplifier (Cairn Research Ltd) digitized using pClamp10 software and a Digidata 1320 A (Molecular Devices). Analysis was performed using Clampfit (Molecular Devices) and Igor software (WaveMeters). Recordings in Syt1−/− and Syt7−/− mice were performed using an Axopatch 200A digitized using IClamp (SciSoft) and a Micro1401 Interface (Cambridge Electronic Design). Recordings in Syt2−/− mice were performed using an EPC-10 patch-clamp amplifier and the Patchmaster software (HEKA). Current recordings were corrected for linear leak conductance measured near ~80 mV. All voltages were corrected for liquid junction potential. For cell capacitance experiments, extracellular solution contained the following (in mM): 115 NaCl, 6 KCl, 5 CaCl2, 1 MgCl2, 30 TEA, 2 Na pyruvate, 8 glucose, 10 Na-HEPES, pH 7.4. Tetrodotoxin (1 μM) and apamin (1 μM) were added to the extracellular solution. Pipette solution for cell membrane capacitance (Cm) experiments contained the following (in mM): 142 CsCl, 1.5 MgCl2, 5 TEA, 1 EGTA, 5 creatine phosphate, 10 Cs-HEPES, pH 7.2. ATP was omitted to minimize activation of purinergic receptors, reported in IHCs after P4 (Trisch and Bergles, 2010). No noticeable difference was observed in changes in cell membrane capacitance (ΔCm) measurements performed in the presence or absence of the nucleotide (data not shown). When EGTA was increased to 5 mM, osmolality was kept constant to 290 mOsm/L by reduction of CsCl. For experiments in Syt2−/− mice, dissections were done in a solution containing the following (in mM): 143 NaCl, 6 KCl, 1.3 CaCl2, 0.9 MgCl2, 0.7 NaH2PO4, 8 glucose, 2 Na-pyruvate, 10 Na-HEPES, pH 7.4. Extracellular recording solution was (in mM): 119 NaCl, 6 KCl, 5 CaCl2, 1 MgCl2, 27 TEA, 2 Na pyruvate, 8 glucose, 10 Na-HEPES, pH 7.4. Tetrodotoxin (1 μM) and apamin (1 μM) were also added to the extracellular solution for Cm measurements. Pipette solution for Cm experiments in Syt2−/− mice contained the following (in mM): 140 Cs-gluconate, 4 MgATP, 0.3 NaGTP, 20 TEA-Cl, 0.5 EGTA, 5 creatine phosphate, 10 HEPES pH 7.2.

For action potential and afferent fiber recordings, the internal solution contained the following (in mM): 140 KCl, 3.5 MgCl2, 0.1 CaCl2, 5 EGTA, HEPES 5, pH 7.2.

Capacitance measurement. Changes in cell membrane capacitance (ΔCm) were used to monitor fusion of synaptic vesicles during exocytosis. Cm was measured according to the Lindau-Neher technique (Lindau and Neher, 1988), using the tracking circuitry of the Optopatch experiments on otoferlin-null mice) or implemented in the IClamp (SciSoft, New Haven, CT) (Syt1-null or Syt7-null mice recordings), or the Patchmaster softwares (Syt2-null mice). A 20 mV amplitude sine wave from a holding potential of ~80 mV was used; for recording of Syt2-null mice, the sine wave had an amplitude of 30 mV from a holding potential of ~90 mV. The resulting maximal depolarizations to ~60 mV were sufficiently small to avoid activation of Ca2+ current (ICa). The command sine wave (781.3–1600 Hz) was blanked during the duration of the voltage step and resumed immediately upon repolarization to capture the capacitance before and after the pulse. ΔCm was estimated as the mean Cm measured 150 μs after the end of the voltage step (to discard capacitance changes due to ICa tail currents) over a period of 50 μs and the prepulse Cm averaged over 50 μs. Only cells with stable series resistance below 10 MΩ (uncompensated) were included in the study.

Afferent fiber recordings. EPSCs were recorded using the whole-cell patch-clamp technique, digitized at 50 kHz, and low pass filtered at 5–10 kHz. Patch pipettes had a resistance of ~10 MΩ.

Single-cell reverse transcription PCR

Inner hair cells were collected from excised organs of Corti (of P1 and P7 mice). Single-cell reverse transcribe (RT)-PCR was done as previously described (Michalski et al., 2007). Synthesis of cDNA was performed using the SuperScript II Reverse Transcriptase kit (Invitrogen). For each cell, we used the extracellular fluid as a negative control. A multiplex nested PCR was performed using specific primers for the tested synaptotagmin (Syt) transcripts (see supplemental Table, available at www.jneurosci.org as supplemental material, for primer sequence) and for two control transcripts, namely myosin VIIa [Myo7A, expressed in both outer hair cells (OHCs) and IHCs] and prestin (only in OHCs) (same primers reported by Michalski et al., 2007). PCR products were sequenced to confirm their specificity. The PCR consisted of 20 amplification cycles (40 s at 95°C, 45 s at 58°C, and 30 s at 72°C). Amplified DNA fragments had a molecular weight ranging between 300 and 500 bp. The nested PCR was performed in a 25 μl mix containing 2 μl of the first PCR mix and only one set of inner primers at a time through 35 cycles (40 s at 95°C, 45 s at 58°C, and 50 s at 72°C).

Twenty IHCs were collected for each developmental stage. Only IHCs that showed the myosin VIIa-positive and prestin-negative expression profiles were taken into account for result analysis (n = 13). The number of Syt-positive cells was then counted for each Syt and compared statistically to the number of extracellular fluid control samples (n = 20) that abnormally scored positive for the presence of cDNA amplification, using the χ2 test (with Yates correction for small samples) to check for significance (p < 0.05).

Whole-mount immunohistochemistry

Cochleae were fixed with 4% paraformaldehyde in PBS for 1 h at 4°C, and the external bone was removed. The organ of Corti was isolated and the tectorial membrane was removed. The tissue samples were permeabilized with 0.3% Triton X-100 in PBS containing 20% normal goat serum for 1 h at room temperature. After PBS washes, they were incubated with the primary antibodies, diluted in PBS containing 10% goat serum (or horse serum) and 0.1% Triton X-100, overnight at 4°C. Omission of the primary antibody was used as a negative control. Primary antibodies were used at the following dilutions: anti-Syt1 (1: 200, rabbit polyclonal, Synaptic Systems; 1:500 mouse monoclonal (C141.1), Synaptic Systems; 1:500 rabbit polyclonal, Abcam), anti-Syt7 (1:200, rabbit polyclonal, Synaptic Systems), anti-Syt2 (znp-1 1:200, mouse monoclonal, ZIRC; or 1:200, rabbit polyclonal against Syt2).
wild-type mice using polyclonal anti-Syt1 (Synaptic Systems) and monoclonal anti-Syt2 (ZIRC) antibodies could be established by the lack of staining in the Syt1−/− or Syt2−/− mutant mice that lack Syt1 and Syt2, respectively (supplemental fig. 2, available at www.jneurosci.org as supplemental material). The same Syt1 immunostaining was observed in the organ of Corti from wild-type mice using any of the three antibodies directed against this protein (data not shown).

Results
Exocytosis in IHCs from early postnatal mice is otoferlin independent
Otoferlin is critical for Ca2+-triggered synaptic exocytosis in IHCs from P6 onwards (Roux et al., 2006). However, IHC exocytosis has not been explored at earlier developmental stages. We thus characterized the Ca2+-dependent exocytosis in IHCs from Otof−/− mice and control littermates from postnatal days 1–8 (P1–P8). To this purpose, we monitored ΔCm in response to voltage activation of Ica.

Electrophysiological data obtained from Otof+/− and Otof−/− IHCs were pooled and referred to as “controls”, after checking that there was no statistical difference between the two groups (unpaired t test, p < 0.01).

In P1 control and Otof−/− IHCs, substantial ΔCm responses could be recorded (Fig. 1A, B). In P1 Otof−/− IHCs, the mean ΔCm evoked by a 100 ms voltage-step to −10 mV (Ica peak) was 17.5 ± 1.7 fF (n = 25), a value similar to that found in control IHCs (14.9 ± 3.1 fF; n = 12; p = 0.12) (Fig. 1C). These ΔCm responses were abolished by 50 μM nifedipine (data not shown, n = 2), indicating that this otoferlin-independent exocytosis is also triggered by L-type Ca2+ channels. The ΔCm responses in Otof−/− IHCs dramatically decreased at P4, and became undetectable from P5 on (Fig. 1C,F), as previously reported (Roux et al., 2006). Notably, a gradual increase of otoferlin immunolabeling in IHCs was observed during the first week of postnatal development: the staining was weak as previously reported (Roux et al., 2006). In P1 Otof−/− IHCs, synaptophysin (1:200 mouse monoclonal, Sigma; 1:200 rabbit polyclonal, Synaptic Systems), anti-CtBP2 (1:200, goat polyclonal, Santa Cruz Biotechnology), anti-o-toferlin (1:500), chicken anti-goat IgG coupled to Alexa Fluor 488 (1:1000). The antibodies for that room temperature. Dilutions of secondary antibodies (1:200 mouse monoclonal, Sigma; 1:200 rabbit polyclonal, Synaptic Systems), anti-CtBP2 (1:200, goat polyclonal, Santa Cruz Biotechnology). Tissue samples were washed in PBS, and incubated with secondary antibodies for 2 h at room temperature. Dilutions of secondary antibodies were as follows: goat anti-rabbit IgG coupled to Alexa Fluor 488 (1:500); goat anti-mouse IgG antibody conjugated with Cy3 fluorophore (Jackson Immunoresearch Laboratories); goat anti-mouse IgG coupled to Alexa Fluor 488 (1:500), chicken anti-goat IgG coupled to Alexa Fluor 488 (1:1000). The nuclear dye DAPI was added to aid in hair cell visualization. The samples were analyzed using a confocal laser scanning microscope LSM510 Meta (Zeiss, Paster Institute, Imagepele) or a Leica confocal upright microscope (Leica DMR TCS SP2 AOBS, Bordeaux Imaging Center). The images were taken with a step size of 0.4 μm. Specificity of anti-Syt antibodies was determined by the lack of residual Syt-immunostaining in the organ of Corti of the respective Syt-null mutant mice. The persistence of the IHCs immunostaining observed in Syt2-null mice when using anti-Syt2 polyclonal antibody (from Synaptic Systems) indicated the nonspecificity of this labeling (data not shown). Therefore, the Syt2 distribution in the organ of Corti was only documented using the znp-1 monoclonal antibody. Similarly, the distribution of Syt2 could not be reliably analyzed since the immunostaining of the organ of Corti obtained with the anti-Syt7 polyclonal antibody (Synaptic Systems) was still detected in Syt7−/− mutant mice (data not shown). In contrast, the specificity of the IHC immunolabeling that was observed in

Syt2, Synaptic Systems), anti-neurofilament NF-200 (1:1000, mouse monoclonal, a gift from Dr. D. Dahl, Harvard Medical School, Boston, MA), anti-synaptophysin (1:200 mouse monoclonal, Sigma; 1:200 rabbit polyclonal, Synaptic Systems), anti-CtBP2 (1:200, goat polyclonal, Santa Cruz Biotechnology). Tissue samples were washed in PBS, and incubated with secondary antibodies for 2 h at room temperature. Dilutions of secondary antibodies were as follows: goat anti-rabbit IgG coupled to Alexa Fluor 488 (1:500); goat anti-mouse IgG antibody conjugated with Cy3 fluorophore (Jackson Immunoresearch Laboratories); goat anti-mouse IgG coupled to Alexa Fluor 488 (1:500), chicken anti-goat IgG coupled to Alexa Fluor 488 (1:1000). The nuclear dye DAPI was added to aid in hair cell visualization. The samples were analyzed using a confocal laser scanning microscope LSM510 Meta (Zeiss, Paster Institute, Imagepele) or a Leica confocal upright microscope (Leica DMR TCS SP2 AOBS, Bordeaux Imaging Center). The images were taken with a step size of 0.4 μm.

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IHC spiking is likely to be at the origin of the EPSCs and spontaneous APs that we recorded in afferent fibers from Otof−/− and control mice, with similar amplitudes and event frequencies (supplemental Fig. 1, available at www.jneurosci.org as supplemental material). These results suggest that exocytosis triggered by spontaneous APs in P1–P3 IHCs does not require otoferlin either.

We further characterized the otoferlin-independent synaptic exocytosis in early postnatal IHCs by studying the synaptic transfer function, that is, the relationship between Ca2+ entry and exocytosis. The voltage-evoked ΔCm responses (100 ms step ranging from −60 to −10 mV) were plotted against the corresponding Ica amplitudes (Fig. 2A,B). The Ca2+ dependence of transmitter release in central synapses, notably at the large calyx of Held nerve terminals, is described by a nonlinear model of cooperative Ca2+ binding to the sensor (Schneckeburger and Neher, 2000; Wölfel and Schneckeburger, 2003). Considering that Ca2+-evoked vesicle fusion (ΔCm) in IHCs is also intrinsically cooperative (Beutner et al., 2001), the data were fitted using the equation ΔCm = g [Ica]n, with Ica varying both as a function of open Ca2+ channels (number increasing with depolarization) and Ca2+ flux per channel (the Ca2+ driving force decreasing with depolarization). In IHCs from both control and Otof−/− P1–P3 mice, the synaptic transfer function displayed a mean power N = 1.7 ± 0.2 (parameters fitted separately to each cell) (Fig. 2B).

Moreover, the otoferlin-independent (P1–P3) and otoferlin-dependent (P7) synaptic transfer functions were similar in the control mice (N = 1.6 ± 0.5 for P7 IHCs, n = 7; p = 0.8) (Fig. 2C). These results suggest that otoferlin-dependent and otoferlin-independent exocytosis in immature IHCs display comparable Ca2+ dependencies, notably with a similar cooperative index.

We compared the dynamics of vesicle pool depletion in IHCs from Otof−/− and control newborn mice (P1) by using a constant depolarizing step (−80 to −10 mV) of duration increasing from 10 to 3000 ms (Fig. 2D–F). Both in control and Otof−/− mice, data points (ΔCm vs time) were best fitted by two exponentials, suggesting the presence of two vesicular pools: a readily releasable pool (RRP) and a slowly releasable pool (SRP) that may arise from vesicles docked at the active zones and vesicles located further from the Ca2+ sensors, respectively. The RRP component of control and Otof−/− P1 IHCs, spanning between 10 and 200 ms, was fitted using a single-exponential with time constants τ = 48 and 63 ms, and ΔCm(max) = 21 fF and 16 fF, respectively (Fig. 2D, inset) (not statistically different, with p = 0.5, in comparing individual cell fit values). Assuming a ΔCm of 37 aF per vesicle (Lenzi et al., 1999), RRP sizes were estimated at 562 and 434 vesicles, respectively. For longer stimuli, control and Otof−/− P1 IHCs displayed similar SRP components (τ = 1363 and 1739 ms, maximal ΔCm = 167 fF and 138 fF, respectively; Fig. 2D).

In P7 Otof−/− IHCs, we confirmed the absence of vesicular release from the RRP (no ΔCm < 500 ms voltage-steps), but found a residual slow exocytosis for longer stimuli (Fig. 2F), as previously reported (Roux et al., 2006).

To address the issue of the relative spatial organization of the Ca2+ channels and Ca2+ sensor, we measured the sensitivity of the RRP and SRP components to various intracellular Ca2+ buffer concentrations. Raising the intracellular concentration of the slow Ca2+ buffer EGTA from 1 to 5 mM, a way to limit intracellular Ca2+ spread to within the microdomains, suppressed the SRP-release but not the RRP-release in IHCs from both control and Otof−/− P1 mice (Fig. 2E). This indicates that Ca2+ channels and non-otoferlin Ca2+ sensors trigger RRP vesicular release work in close vicinity in P1 IHCs. This result also suggests that SRP release results from vesicles located farther from the Ca2+ channels, or, alternatively, might reflect a Ca2+-dependent recovery of the RRP. Furthermore, repetitive exocytosis of the RRP (evoked by a train of 50 ms depolarizing...
steps) was similar in control (see Fig. 4G) and Otof−/− P1 IHCs (data not shown), with an average cumulative \( C_{\text{cm}} \) response of 21.8 ± 0.5 fF \((n = 6)\) and 18.4 ± 4.1 fF \((n = 4)\), respectively, which suggests that the recruitment of synaptic vesicles to maintain a constant RRP in P1 IHCs is not affected in the absence of otoferlin. In summary, all parameters of exocytosis tested in P1–P2 Otof−/− mice were unchanged compared with their control littermates.

**Expression of synaptotagmins in IHCs**

The persistence of Ca\(^{2+}\)-dependent exocytosis in IHCs from P0–P3 Otof−/− mice raises the question of the molecular identity of the Ca\(^{2+}\) sensors involved at early postnatal stages. Since several Syts are regarded as Ca\(^{2+}\) sensors regulating exocytosis in the CNS, we used single-cell RT-PCR to detect Syt transcripts in IHCs. We investigated the expression of eight different Syts, Syt1, 2, 3, 5, 6, 7, 9, and 10, all known for their Ca\(^{2+}\)-binding properties (Su¨dhof, 2002), in P1 and P7 IHCs. Syt1, 2, 6, and 7 displayed the highest expression levels in P1 IHCs, while Syt2 was markedly decreased (Fig. 3G). These results were further substantiated by immunolabeling experiments and confocal imaging of the organ of Corti before (P1, P6, P8) and after (P15 and P21) the onset of hearing. Whole-mount preparations were double-immunostained for Syt1 or Syt2, and for the synaptic vesicle protein synaptophysin. This latter protein is a marker of lateral efferent presynaptic terminals, that form axodendritic synapses with afferent dendrites below IHCs, and medial efferent synapses contacting OHCs (Gil...
Loyzaga and Pujol, 1988; Safieddine and Wenthold, 1999). From P1 to P6, Syt1 and Syt2 labelings were present in both IHCs and OHCs (Fig. 3A–C), but the two proteins displayed different subcellular distributions as shown by double-labeling experiments (Fig. 3B, C). The strongest Syt1-immunolabeling was observed along the basolateral membrane that lines the IHC presynaptic region (Fig. 3A, B), whereas the Syt2-immunolabeling was mainly cytoplasmic (Fig. 3B, C). In addition, Syt1 and Syt2 displayed different temporal profiles in the IHCs, as expected from our single-cell RT-PCR results. The Syt2-immunolabeling was no longer observed in IHCs from P8 onward, while Syt1 could still be detected up to P10 through all the cochlear spiral (Fig. 3D; supplemental Fig. 3, available at www.jneurosci.org as supplemental material), but vanished at P15 in the IHCs of the cochlear middle turn (Fig. 3E, F) and at P21 in those of the apical turn (data not shown). Finally, at all stages investigated (from P1 to P21), Syt1 and Syt2 immunolabelings were detected in the efferent, synapticysphin-inmunoreactive, nerve fibers (Fig. 3A–F; data not shown).

In Otof−/− IHCs, Syt1 and Syt2 had the same distributions and developmental patterns, as in control littermates (data not shown). Furthermore, Syt2 was present in P0 Syt1−/− IHCs and Syt1 was present in P0 and P6 Syt2−/− IHCs (data not shown), suggesting that the lack of one Ca2+ sensor does not influence the expression of the other.

Syt1−/− IHCs exhibit a RRP release unchanged but a diminished RRP recovery

We next assessed the role of Syt1 in IHC exocytosis by measuring Cm changes in response to ICa in P0 Syt1−/− mice. The short postnatal survival of these mice (~24 h) (Geppert et al., 1994) restricted measurements to newborn mice. No morphological anomalies were observed in the IHCs and OHCs of the Syt1−/− newborn mice. Indeed, we found presynaptic ribbons and proper innervation contacting the hair cells, with no decrease in the number of cochlear ganglion neurons (Fig. 4A).

In Syt1−/− P0 IHCs, ICa-evoked ΔCm responses were similar to those in Syt1+/− (wild-type) IHCs (Fig. 4B, C), with a mean amplitude of 25.6 ± 6.1 fF (n = 11) and 28.5 ± 8.8 fF (n = 11), respectively (p = 0.9) for a 100 ms voltage-step close to ICa peak. The corresponding ICa amplitudes were 87 ± 9 pA and 86 ± 9 pA (p = 0.8) (Fig. 4C). The synaptic transfer function in Syt1−/− IHCs was evaluated as described above, by plotting ΔCm against ICa (100 ms voltage-step ranging from −60 mV to −10 mV, from a holding potential of −80 mV) (Fig. 4D). In wild-type and Syt1−/− IHCs, the synaptic transfer function was best fitted using N = 2.1 ± 0.6 (n = 11) and N = 2.0 ± 0.2 (n = 10), respectively (p = 0.4), indicating that exocytosis evoked in the absence of Syt1 has an unchanged cooperative index in P0 IHCs. The kinetics of RRP exocytosis (~10 mV voltage-steps of various durations) were also similar in Syt1−/− mice and control littermates (τ = 60 ± 20 ms and 65 ± 14 ms, respectively; p = 0.8; Fig. 4E). This suggested that release from the RRP was unchanged in newborn Syt1−/− mice.

We observed, however, that depolarizing stimuli had to be separated by at least 1 min intervals to trigger reproducible ΔCm responses in Syt1−/− IHCs. This long time requirement after each Ca2+ stimulation suggested that the recovery of RRP vesicles might be affected in Syt1−/− IHCs. To further investigate the vesicle recruitment and refilling of the RRP, we stimulated IHCs with a train of brief (50 ms) voltage steps spaced by 100 ms at the holding potential of −80 mV, during which sine wave for Cm measurements was applied (Fig. 4F, top). In wild-type mice, a 50 ms-depolarizing train evoked a linear increase of the cumulative ΔCm values (Fig. 4F, G), indicating that the RRP was able to be rapidly refilled in-between successive stimuli. In contrast, in Syt1−/− IHCs, the RRP recruitment was affected after the second test pulse, despite a stable ICa amplitude (Fig. 4F, G). The ΔCm at
Ca$^{2+}$-dependent exocytosis is not affected in IHCs from Syt2$^{-/-}$ or Syt7$^{-/-}$ mice

Syt2$^{-/-}$ mice generally survive up to 3 weeks, but exhibit severe motor dysfunction after the second week (Pang et al., 2006). Given the selective expression of Syt2 in IHCs during the first few postnatal days (Fig. 3), we first studied exocytosis in P2–P3 Syt2$^{-/-}$ IHCs (Fig. 5), but we failed to detect deficits in exocytosis. In Syt2$^{-/-}$ or Syt2$^{-/-}$ control mice, a 100 ms voltage step to −10 mV evoked a Ca$^{2+}$ current ($I_{\text{Ca}}$) of 172 ± 19 pA, which elicited a Δ$C_m$ step of 6.8 ± 1.3 fF (n = 17). In Syt2$^{-/-}$ IHCs, a same voltage stimuli elicited an average Δ$C_m$ response of 6.3 ± 1.3 fF (n = 10), which was indistinguishable from the one measured in the control mice (p > 0.5). The Ca$^{2+}$ current was 125 ± 23 pA (n = 10); which was slightly, but not significantly smaller than the one measured in control mice (p = 0.13). In plots of the exocytotic Δ$C_m$ response versus the length of the depolarizing pulses, the release kinetics was not substantially modified in Syt2$^{-/-}$ mice compared with control littermates (Fig. 5C). Using a train of brief (50 ms) voltage steps to −10 mV (close to peak of $I_{\text{Ca}}$), a sustained vesicular release, similar to controls, could be observed in IHCs from P2–P3 Syt2$^{-/-}$ mice, indicating that the RRP was rapidly refilled between voltage pulses (Fig. 5D).

Since Syt2 has been recently proposed to underlie the Ca$^{2+}$ dependence of vesicle release in mature IHCs (Johnson et al., 2010), we also investigated the properties of exocytosis in near-mature, P15–P17 IHCs from Syt2$^{-/-}$ mice (Fig. 6). Again, we did not detect any obvious exocytotic defect. In Syt2$^{-/-}$ P15–P17 IHCs, 100 ms voltage steps to $I_{\text{Ca}}$ peak evoked Δ$C_m$ with a mean amplitude of 27.1 ± 2.3 fF for $I_{\text{Ca}}$ peak of 183 ± 11 pA (n = 15). These values were similar to those obtained in the wild-type littermates [27.4 ± 4.7 fF and 182 ± 23 pA, n = 8 (p > 0.5 for both comparisons)]. In P15–P17 IHCs from Syt2$^{-/-}$ mice, both the kinetics of exocytosis (Fig. 6C) and vesicle recruitment (Fig. 6D) were similar to those in the control littermates. Furthermore, the
synaptic transfer function showed similar linear relationships in P15–P17 wild-type and Syt2+/−/− mice (Fig. 6E).

In Syt7+/−/− mice, we did not find any defect of Ca2+-evoked exocytosis (RRP component) induced by \( I_{\text{Ca}} \) activation in immature (P2–P8) or post-hearing onset (P15–P30) IHCs (Table 1, Fig. 7). In these mice, Ca2+-evoked exocytosis underwent a normal postnatal maturation, with an increase in Ca2+ efficiency and a linearization of its Ca2+ dependency after hearing onset. A train of short depolarizations was also able to induce sustained vesicular release, which is indicative of a normal RRP refilling (data not shown).

Discussion

The here uncovered switch from an otoferlin-independent to an otoferlin-dependent exocytosis, reveals an as yet unknown maturation step of the IHC synaptic apparatus that takes place around P4 in the mouse. At this developmental transition, the spontaneous Ca2+-driven spiking activity of IHCs rises to its highest frequency (Marcotti et al., 2003), and is generated both intrinsically and extrinsically through ATP-dependent excitation of IHCs (Tritsch and Bergles, 2010). The rise in IHC spike frequency and resulting increase in glutamate release, are supposed to be important for IHC synaptic maturation (Seal et al., 2008) and for proper maturation of the central auditory pathway (Rubel and Fritzsch, 2002; Leake et al., 2006; Leao et al., 2006). Notably, this period of postnatal cochlear development corresponds to an extensive period of synaptic reorganization, refinement and retraction of type I and type II afferent neurites contacting IHCs (Lenoir et al., 1980; Huang et al., 2007). Our results show that before P4, immature IHCs make use of other Ca2+ sensors than otoferlin to drive Ca2+-dependent exocytosis. In that respect, it is noteworthy that a \( \Delta C_m \) response has been recorded in IHCs from hypothyroid rats, in which otoferlin could not be detected (Brandt et al., 2007; Sendin et al., 2007). The otoferlin-independent exocytosis phase reported here may account for this observation. Indeed, it has been shown that the maturation of the auditory neuroepithelium is delayed in thyroid hormone-deficient mice (Sendin et al., 2007).

Before P3, a period during which IHC exocytosis is otoferlin-independent, Syt1, Syt2 or Syt7 were found to be dispensable for vesicle fusion, indicating that other, as yet unknown Ca2+ sensors, are involved in IHCs. Interestingly, we found that in response to repetitive stimulation, IHC exocytosis was seriously affected in Syt1−/− mice, suggesting that Syt1 is involved in the recruitment of synaptic vesicles to the RRP, possibly by ensuring fast Ca2+-dependent vesicle replenishment of the ribbon synapse. In CNS nerve terminals, Syt1 is also known to regulate the rate of endocytosis of synaptic vesicles that have undergone fusion (Poskanzer et al., 2003; Nicholson-Tomishima and Ryan, 2004). The replenishment defect of the RRP observed in Syt1−/− IHCs could therefore result from an impaired clearance of recently exocytosed vesicles from release sites (Hosoi et al., 2009).

The absence of Syt1 immunolabelling in IHCs after the onset of hearing, however, suggests that this Ca2+ sensor is not involved in mature IHCs. Furthermore, post-hearing-onset IHCs from Syt2−/− or Syt7−/− mice display normal exocytosis, indicating that Syt2 and Syt7 are also not essential for exocytosis in mature IHCs. This result is consistent with the absence of Syt2 in IHCs beyond the first postnatal week. Since several Syt isoforms are expressed in immature IHCs, we cannot exclude at present that the lack of a strong phenotype in single Syt-null mice may be due to functional redundancy between Syts, although we note that a redundancy between Syt isoforms has not been shown so far in other synapses. Complexins, that are also known to bind to the SNARE complex and regulate fast exocytosis in conjunction with Syts in most neurosecretory synapses, are also dispensable at the mature IHC ribbon synapse (Strenzke et al., 2009). Together, these results suggest that the mature IHC makes use of an unconventional synaptic machinery, which includes otoferlin, to control exocytosis. The recent report of a temperature-sensitive form of deafness caused by an otoferlin mutation, that manifests a sudden hearing impairment provoked by fever in otherwise normal hearing children, further argues in favor of otoferlin playing a critical role at the synapses of mature IHCs (Marlin et al., 2010).

Surprisingly, exocytosis of P7 immature IHCs, while otoferlin-dependent, still showed a low Ca2+ efficiency compared with mature IHCs (Fig. 7). Indeed, the Ca2+ threshold (minimum \( I_{\text{Ca}} \) required to evoke a release of vesicles >5 FF) was similar in P1 IHCs (otoferlin not required) and P7 IHCs (otoferlin required), while it is much lower in post-hearing onset (P15–

### Table 1. Ca2+-dependent exocytosis is not affected in Syt7−/− IHCs

<table>
<thead>
<tr>
<th>Age</th>
<th>Genotype</th>
<th>( \Delta C_m (\text{max}) (\text{fF}) )</th>
<th>( \Delta C_m/I_{\text{Ca}} (\text{fF} \text{pA}^{-1}) )</th>
<th>Ca2+ dependency Npower</th>
</tr>
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<tbody>
<tr>
<td>P2–P3</td>
<td>Syt7+/−/−</td>
<td>12.5 ± 1.9 (n=3)</td>
<td>0.13 ± 0.02 (n=3)</td>
<td>–</td>
</tr>
<tr>
<td>Syt7+/−</td>
<td>11 ± 0.2 (n=3)</td>
<td>0.12 ± 0.02 (n=3)</td>
<td>2.1 ± 0.2 (n=2)</td>
<td></td>
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<tr>
<td>P6–P8</td>
<td>Syt7+/+</td>
<td>20 ± 4 (n=6)</td>
<td>0.12 ± 0.02 (n=6)</td>
<td>2.2 ± 0.2 (n=4)</td>
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<tr>
<td>P15–P30</td>
<td>Syt7+/+</td>
<td>24.3 ± 6 (n=8)</td>
<td>0.17 ± 0.04 (n=8)</td>
<td>2.2 ± 0.1 (n=6)</td>
</tr>
<tr>
<td>Syt7+/−/−</td>
<td>66 ± 11 (n=6)</td>
<td>0.28 ± 0.05 (n=6)</td>
<td>0.9 ± 0.1 (n=5)</td>
<td></td>
</tr>
<tr>
<td>Syt7+/−/−</td>
<td>69 ± 11 (n=5)</td>
<td>0.37 ± 0.08 (n=5)</td>
<td>1.1 ± 0.2 (n=5)</td>
<td></td>
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Figure 7. Comparative Ca2+ thresholds triggering exocytosis in IHCs during postnatal development. A, Comparative IHC synaptic transfer functions (\( \Delta C_m (\text{max}) \) vs \( I_{\text{Ca}} \)) plotted on a double log scale, for P1–Otof−/− (open triangle), P1–Otof+/− (black circle) (data from Fig. 2B), P0–Syt1−/− (gray circle; data from Fig. 4D), P7–Otof−/− (green triangle), adult (P15–P30) Syt7−/− (open square) and adult (P16–P30) Syt7−/− (blue square) IHCs. The \( I_{\text{Ca}} \) thresholds necessary to evoke significant \( \Delta C_m (>5 \text{FF}) \) were extrapolated by fitting (line) the slope of the initial \( C_m \) increase. B, Synthetic vesicular release in adult IHCs has a higher Ca2+ sensitivity (4– to 5-fold) than in P0–P1 and P7 IHCs. This increase in Ca2+ sensitivity in mature IHCs is most likely not due to otoferlin on its own, since P7 IHCs, which still have low Ca2+ sensitivity of exocytosis, already require otoferlin for synaptic vesicle release.
Lacking to set a linear, highly Ca\textsuperscript{2+} sensitive, synaptic function in P6 IHCs. Synaptotagmin 4, present in mature IHCs only, might be one of the otoferlin partners, since it has been shown to be required for the linearization of the synaptic transfer function (Johnson et al., 2010). A different distribution of Ca\textsuperscript{2+} channels at the synaptic active zones between immature and mature IHCs is also expected to change the synaptic transfer function. In P6 IHCs, Ca\textsuperscript{2+} channels are not exclusively located at the active zone (Zampini et al., 2010), and the extrasynaptic location of some Ca\textsuperscript{2+} channels may contribute to the lower “Ca\textsuperscript{2+} efficiency” of exocytosis in immature IHCs.

Otoferlin is a member of the ferlin protein family, which includes fer-1, dysferlin and myoferlin. When defective, dysferlin and myoferlin are responsible for muscular dystrophies (Bansal and Campbell, 2004). In Caenorhabditis elegans, Fer-1 is required for Ca\textsuperscript{2+}-dependent vesicle fusion in spermatids during spermiogenesis (Achanzar and Ward, 1997; Washington and Ward, 2006). Based on the muscular phenotypes of the dysferlin and myoferlin knock-out mice, a role of these proteins in Ca\textsuperscript{2+}-dependent membrane repair processes and myoblast membrane fusion, respectively, has been proposed (Bansal et al., 2003; Doherty et al., 2005). Otoferlin, whose C2 domains bind to SNARE proteins in a Ca\textsuperscript{2+}-dependent manner (Roux et al., 2006), is critically involved in membrane vesicle fusion in IHCs from ~P4–P5 onwards. In an in vitro fusion assay, otoferlin accelerates membrane fusion with higher Ca\textsuperscript{2+} affinity than Syt1 (C. Johnson and E. R. Chapman personal communication). In vestibular hair cells, otoferlin has been shown to be essential for a high affinity Ca\textsuperscript{2+} sensor function that allows fast exocytosis during brief cell depolarizations and low Ca\textsuperscript{2+} stimuli (Dulon et al., 2009). The requirement for otoferlin at the mature IHC synapse is likely due to a constraint specific to the auditory ribbon synapse organization, and/or its mode of exocytosis that involves multivesicular release (Glowatzki and Fuchs, 2002). It is possible that the larger number of C2 domains in otoferlin than in Syt1 (six vs two) enables more complex interactions between otoferlin and SNARE proteins, perhaps controlling the supply of vesicles to the release sites, as recently suggested in pachanga mice (Pangrscik et al., 2010) and/or allowing a variety of vesicle fusion modes. In the mature IHC, which does not contain Syt1 or Syt2, the proportion of coordinated multivesicular fusion events at the ribbon synapse has been shown much higher than in Syts-expressing IHCs from pre-hearing animals (Grant et al., 2010). We thus suggest that otoferlin has a synchronizing effect on the multivesicular fusion events, an effect that could be attenuated by the coexistence of Syts, notably Syt1, in immature IHCs.

References


