

Bridging the gap between presynaptic hair cell function and neural sound encoding

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Abstract

Neural diversity can expand the encoding capacity of a circuitry. A striking example of diverse structure and function is presented by the afferent synapses between inner hair cells (IHCs) and spiral ganglion neurons (SGNs) in the cochlea. Presynaptic active zones at the pillar IHC side activate at lower IHC potentials than those of the modiolar side that have more presynaptic Ca^{2+} -channels. The postsynaptic SGNs differ in their spontaneous firing rates, sound thresholds and operating ranges. While a causal relationship between synaptic heterogeneity and neural response diversity seems likely, experimental evidence linking synaptic and SGN physiology has remained difficult to obtain. Here, we aimed at bridging this gap by *ex vivo* paired recordings of IHCs and postsynaptic SGN boutons with stimuli and conditions aimed to mimic those of *in vivo* SGN-characterization. Synapses with high spontaneous rate of release (SR) were found predominantly on the pillar side of the IHC. These high SR synapses had larger and more compact spontaneous EPSCs, lower voltage-thresholds, tighter coupling of Ca^{2+} channels and vesicular release sites, shorter response latencies and higher initial release rates. This study indicates that synaptic heterogeneity in IHCs directly contributes to the diversity of spontaneous and sound-evoked firing of SGNs.

Significance Statement

Sound encoding relies on spiral ganglion neurons (SGNs) with diverse spontaneous firing, sound thresholds of firing and sound-intensity range over which SGN firing rate changes. Such functional SGN diversity might originate from different input from afferent synapses with inner hair cells (IHCs). The present study addresses this hypothesis by using recordings from individual IHC-SGN synapses of hearing mice under *ex vivo* conditions aimed to mimic cochlear physiology. The results provide evidence that synaptic heterogeneity in IHCs contributes to SGN firing diversity. Thus, the cochlea employs heterogeneous synapses to decompose sound information into different neural pathways that collectively inform the brain about sound intensity.

eLife assessment

This **fundamental** study advances substantially our understanding of sound encoding at synapses between single inner hair cells of the mouse cochlea and spiral ganglion neurons. Dual patch-clamp recordings—a technical tour-de-force—and careful data analysis provide **compelling** evidence that the functional heterogeneity of these synapses contributes to the diversity of spontaneous and sound-evoked firing by the neurons. The work will be of broad interest to scientists in the field of auditory neuroscience.

Introduction

Chemical synapses represent diverse and plastic neural contacts that are adapted to the specific needs of neural computation. Synaptic diversity is expressed across the nervous system, within a given circuit and even within the same neuron (recent reviews in Ref. [Nusser, 2018](#); [Wichmann and Kuner, 2022](#)). Synaptic diversity occurs at various levels: from synapse shape and size, to the ultrastructure of pre- and postsynaptic specializations, to their molecular composition. The auditory system harbors striking examples of synaptic diversity. Glutamatergic ribbon synapses in the cochlea, calyceal synapses in the brainstem, and bouton synapses throughout the central auditory system differ greatly from each other ([Moser et al., 2006](#); [Wichmann and Kuner, 2022](#)). Beyond the diversity across synapses formed by different neurons, heterogeneity exists even among auditory synapses of an individual presynaptic inner hair cell (IHC) with its 5–30 postsynaptic spiral ganglion neurons (SGNs, ([Gómez-Casati and Goutman, 2021](#); reviews in: [Meyer and Moser, 2010](#); [Moser, 2020](#); [Reijntjes and Pyott, 2016](#))). Such synaptic heterogeneity is a candidate mechanism for how the cochlea decomposes acoustic information ([Grant et al., 2010](#); [Meyer et al., 2009](#); [Ohn et al., 2016](#); [Özçete and Moser, 2021](#)). For example, the cochlea might use heterogeneous afferent synapses to break down sound intensity information into complementary spike rate codes of SGNs that have been reported across species ([Huet et al., 2016](#); [Kiang et al., 1965](#); [Sachs and Abbas, 1974](#); [Taberner and Liberman, 2005](#); [Winter et al., 1990](#)).

Decades of *in vivo* recordings from single SGNs have demonstrated functional diversity of SGNs with comparable frequency tuning, i.e., receiving input from IHCs at a given tonotopic place or potentially even the same IHC. Such functional diversity is present in both spontaneous and sound-evoked firing. The spontaneous firing rate (SR) in the absence of sound varies from less than 1 spikes/s to more than 100 spikes/s ([Barbary, 1991](#); [Evans, 1972](#); [Kiang et al., 1965](#); [Schmiedt, 1989](#); [Taberner and Liberman, 2005](#)). In response to increasing sound pressure, SGNs with high SR show a low sound threshold and a steep rise in the spike rate to increasing sound intensities until the rate saturates. SGNs with low SR have a higher sound thresholds, shallower spike rate rise and saturate at higher sound intensities ([Ohlemiller et al., 1991](#); [Winter et al., 1990](#)). Additionally, SGNs show differences in their excitability ([Crozier and Davis, 2014](#); [Markowitz and Kalluri, 2020](#); [Smith et al., 2015](#)), morphological features ([Liberman, 1980](#); [Merchan-Perez and Liberman, 1996](#); [Tsuji and Liberman, 1997](#)) and heterogeneous molecular profiles ([Li et al., 2020](#); [Petitpré et al., 2020](#), 2018; [Shrestha et al., 2018](#); [Sun et al., 2018](#)). Yet, it has remained challenging to demonstrate a causal link of a candidate mechanism to the physiological SGN diversity.

One common approach has been to capitalize on a pioneering study that employed *in vivo* labelling of physiologically characterized SGNs in cats and proposed that synapses formed by low and high SR SGNs segregate on the basal IHC pole ([Liberman, 1982](#)). High SR SGNs preferentially

contacted the pillar side of the IHC (facing pillar cells), while low SR SGNs synapsed on the opposite, modiolar side of the IHC (facing the cochlear modiolus). Interestingly, a segregation has also been found for afferent and efferent synaptic properties, as well as molecular and biophysical SGN properties (Frank et al., 2009 [↗](#); Grant et al., 2010 [↗](#); Hua et al., 2021 [↗](#); Kantardzhieva et al., 2013 [↗](#); Liberman et al., 2011 [↗](#); Markowitz and Kalluri, 2020 [↗](#); Merchán-Pérez and Liberman, 1996 [↗](#); Meyer et al., 2009 [↗](#); Neef et al., 2018 [↗](#); Ohn et al., 2016 [↗](#); Özçete and Moser, 2021 [↗](#); Shrestha et al., 2018 [↗](#); Sun et al., 2018 [↗](#); Yin et al., 2014 [↗](#)). For instance, type I_b and I_c SGNs preferentially synapse on the modiolar side (Sherrill et al., 2019 [↗](#); Shrestha et al., 2018 [↗](#); Sun et al., 2018 [↗](#)) and show low SR (Siebald et al., 2023 [↗](#)). Pillar synapses are preferentially formed by type I_a SGNs (Shrestha et al., 2018 [↗](#); Siebald et al., 2023 [↗](#)), have smaller IHC active zones (AZs) (Liberman et al., 2011 [↗](#); Ohn et al., 2016 [↗](#); Reijntjes et al., 2020 [↗](#)), and activate at voltages as low as the IHC resting potential (Ohn et al., 2016 [↗](#); Özçete and Moser, 2021 [↗](#)). The low voltage of activation of release at pillar synapses shown *ex vivo* could underly the high SR and low sound threshold of firing of SGNs found *in vivo* but direct demonstration of such a link has been missing.

Here, we aimed to bridge the gap between *ex vivo* presynaptic physiology and *in vivo* SGN neurophysiology. We performed paired IHC and SGN bouton recordings in acutely explanted organs of Corti from hearing mice with stimuli and conditions aimed to mimic *in vivo* SGN-characterization. This approach tightly controls IHC Ca^{2+} influx and records the postsynaptic SGN response to glutamate release at a single afferent synapse. Using the rate of spontaneous excitatory postsynaptic currents (rate of sEPSCs, SR) as a surrogate for SGN SR, we demonstrate that high SR synapses have larger and more compact sEPSCs as well as lower voltage thresholds, shorter latencies of evoked EPSCs, tighter Ca^{2+} channel coupling to vesicle release and higher initial release rates. 90% of these high SR synapses were located on the pillar side of the IHC. Our findings suggest that synaptic heterogeneity accounts for much of the SGN firing diversity.

Results

Simultaneous paired patch-clamp recordings from IHCs and one of the postsynaptic SGN boutons were performed on mice after the onset of hearing (postnatal days (p) 14–23). We performed perforated-patch whole-cell configuration from IHCs, held at their presumed physiological resting potential (−58 mV (Johnson, 2015 [↗](#))), and ruptured-patch whole-cell recordings from one of the postsynaptic SGN boutons (Fig. 1 [↗](#)). Due to the technical difficulty of establishing the paired recording, typically only one bouton was recorded per IHC. Recordings were made at body temperature and in artificial perilymph-like solution (Wangemann and Schacht, 1996 [↗](#)). To establish the paired recording, we approached boutons facing either the pillar or the modiolar side of the IHC in an effort to elucidate synaptic differences between both sides (Fig. 1A [↗](#)). We nickname the synapse location as “pillar” and “modiolar” based on the DIC-image, but note that efforts to stain and image the recorded boutons by fluorescence microscopy were not routinely successful. In addition, the recorded boutons were classified based on their spontaneous rate of synaptic transmission (SR, **Low SR** < 1 sEPSC/s vs **High SR** ≥ 1 sEPSC/s according to (Taberner and Liberman, 2005 [↗](#))).

Spontaneous synaptic transmission

In order to recapitulate synaptic transmission in the absence of sound stimulation, we held the IHC at their presumed physiological resting potential (−58 mV (Johnson, 2015 [↗](#)). Since $Ca_v1.3$ Ca^{2+} -channels activate at low voltages (−65 to −45 mV (Koschak et al., 2001 [↗](#); Picher et al., 2017 [↗](#); Platzer et al., 2000 [↗](#); Xu and Lipscombe, 2001 [↗](#))), their open probability at the IHC resting potential is thought to be sufficient to trigger spontaneous release (Glowatzki and Fuchs, 2002 [↗](#); Özçete and Moser, 2021 [↗](#); Robertson and Paki, 2002 [↗](#)). Under our experimental conditions, spontaneous, i.e., excitatory postsynaptic currents in the absence of IHC stimulation (sEPSCs) were

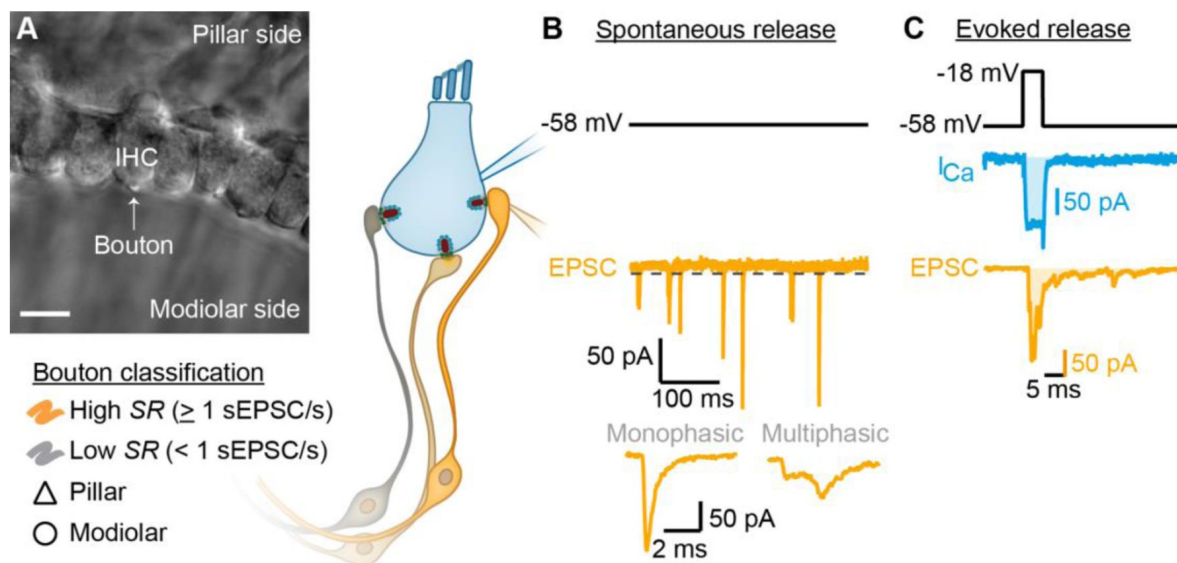


Figure 1.

Paired IHC-bouton patch-clamp recordings to study the release properties of individual IHC ribbon synapses as a function of synapse position

A. Differential Interference Contrast image of an explanted murine organ of Corti. In this example, supporting cells from the modiolar side were removed to gain access to the IHCs and their contacting boutons. The recorded boutons were classified based on their position (Δ pillar or $\circ$ modiolar) and on their spontaneous rate (SR) (**Low SR** < 1 sEPSC/s vs **High SR** ≥ 1 sEPSC/s). Scale bar: 10 μ m. **B.** Spontaneous release was recorded in absence of stimulation (i.e., IHC holding potential = -58 mV; **Supplementary Table 1**; dashed line represents the threshold for sEPSC detection). sEPSCs were classified as monophasic (or compact: a steady rise to peak and monoexponential decay) or as multiphasic (or non-compact: multiple inflections and slowed raising and decaying kinetics). **C.** Evoked release: depolarizing pulses (black trace) were used to trigger whole IHC Ca^{2+} influx (I_{Ca} , blue trace) and ensuing release of neurotransmitter that evoked eEPSCs, light orange trace). Ca^{2+} charge and eEPSC charge were estimated by taking the integral of the currents (shaded light blue and light orange areas).

observed in 23 of 33 pairs (**Fig. 2B**). We used the rate of sEPSCs (*SR*) as a surrogate of SGN *SR* as sEPSCs trigger SGN firing with >90% probability (Rutherford et al., 2012). Regardless of all efforts to maintain the physiological integrity in the *ex vivo* experiments, we expect our estimated rates of sEPSC to underestimate the SGN *SR* for the same age group (Wong et al., 2013). For each paired recording, we quantified the *SR* during 5 or 10 s of a continuous recording, or during the segment before and after the step depolarization protocols (**Supplementary Table 1**). Amplitudes of sEPSCs typically varied from around -10 pA to -400 pA across all recorded IHC-SGN synapses (**Fig. 2B**). The amplitude histogram for all pairs was slightly skewed towards larger amplitudes (skewness of 1.06) with a coefficient of variation (CV) of 0.68. The charge distribution for all pairs displayed a prominent peak at 40 fC, with a skewness of 2.00 and a CV of 0.77. *SR* ranged from 0 to about 18 sEPSC/s (**Fig. 2H**). The *SR* distribution was highly skewed, with a median of 0.2 sEPSC/s. Following the spontaneous firing rate classification of SGNs by Taberner and Liberman (2005), we classified the synapses into low (< 1 sEPSC/s, ~70%) or high (≥ 1 sEPSC/s, ~30%) *SR* synapses. Next, we analyzed the recordings as a function of i) synapse position and ii) rate of spontaneous synaptic transmission.

i) Position dependence of synaptic transmission

Of the 33 obtained paired recordings, 20 were classified as pillar synapses. *SR* was significantly higher for the pillar synapses (mean *SR* of 2.30 ± 0.96 sEPSC/s; median 0.59; $n = 20$) compared to modiolar ones (mean *SR* of 0.22 ± 0.09 sEPSC/s; median 0.05; $n = 13$; $p = 0.0311$, Mann-Whitney U test). The mean amplitude (**Fig. 2C**) and charge (**Fig. 2D**) of the sEPSCs were comparable between modiolar and pillar SGN boutons (Amp_{sEPSC} -69.37 ± 13.91 pA ($n = 8$ modiolar) vs -87.21 ± 9.49 pA ($n = 13$ pillar); $p = 0.2913$, unpaired t-test; Q_{sEPSC} 63.74 ± 7.77 fC (modiolar) vs 72.54 ± 9.57 fC (pillar); $p = 0.5463$, unpaired t-test). sEPSC of pillar SGN boutons showed significantly shorter 10-90% rise times than the modiolar ones (**Figure 2-figure supplement 1**, panel A and C; 0.38 ± 0.04 ms vs 0.57 ± 0.06 ms; $p = 0.0111$, unpaired t-test), yet similar decay times and full-width half-maximum (FWHM, **Figure 2-figure supplement 1**, panels A, D-E; $p = 0.7997$ and $p = 0.9198$, respectively, unpaired t-test). As a second approach to sEPSC properties, we quantified the percentage of monophasic (or compact) sEPSCs (Chapochnikov et al., 2014; Glowatzki and Fuchs, 2002) and found a non-significant trend towards higher percentages of monophasic sEPSCs for pillar synapses (**Fig. 2G**; $25.57 \pm 6.9\%$ for modiolar vs $46.65 \pm 7.06\%$ for pillar; $p = 0.0681$, unpaired t-test).

ii) Relation of synaptic properties to the rate of spontaneous synaptic transmission

High *SR* synapses had significantly larger sEPSCs (**Fig. 2J** and **Figure 2-figure supplement 1**, panel Bi; average sEPSC amplitude of -105.2 ± 8.47 pA for high *SR* ($n = 10$) vs. -62.39 ± 9.67 pA for low *SR* ($n = 13$); $p = 0.0042$, unpaired t-test). Likewise, sEPSC charge tended to be larger in high *SR* SGN synapses (**Fig. 2K** and **Figure 2-figure supplement 1**, panel Bii; Q_{sEPSC} 84.23 ± 10.38 fC for high *SR* vs. 58.14 ± 7.79 fC for low *SR*; $p = 0.0527$, unpaired t-test). The fraction of monophasic sEPSCs was significantly higher in high *SR* synapses (**Fig. 2L**; $52.08 \pm 6.65\%$; median 43.71%) compared to low *SR* synapses ($29.49 \pm 7.43\%$; median 22.22%; $p = 0.0185$, Mann-Whitney U test). High *SR* synapses also showed a significantly faster 10-90% rise times (**Figure 2-figure supplement 1**, panel F; 0.36 ± 0.04 ms) than low *SR* synapses (0.51 ± 0.05 ms; $p = 0.0420$, unpaired t-test).

Other sEPSCs kinetics, such as decay time constant and FWHM, were not different between low and high *SR* pairs (**Figure 2-figure supplement 1**, panel G,H; $p = 0.7969$ and $p = 0.9948$, respectively, unpaired t-test). Taken together, these results indicate that high *SR* synapses are characterized by sEPSCs with larger amplitudes and charges, faster rising times and a more compact waveform. Interestingly, 9 out of 10 high *SR* synapses were located on the pillar side of the IHC.

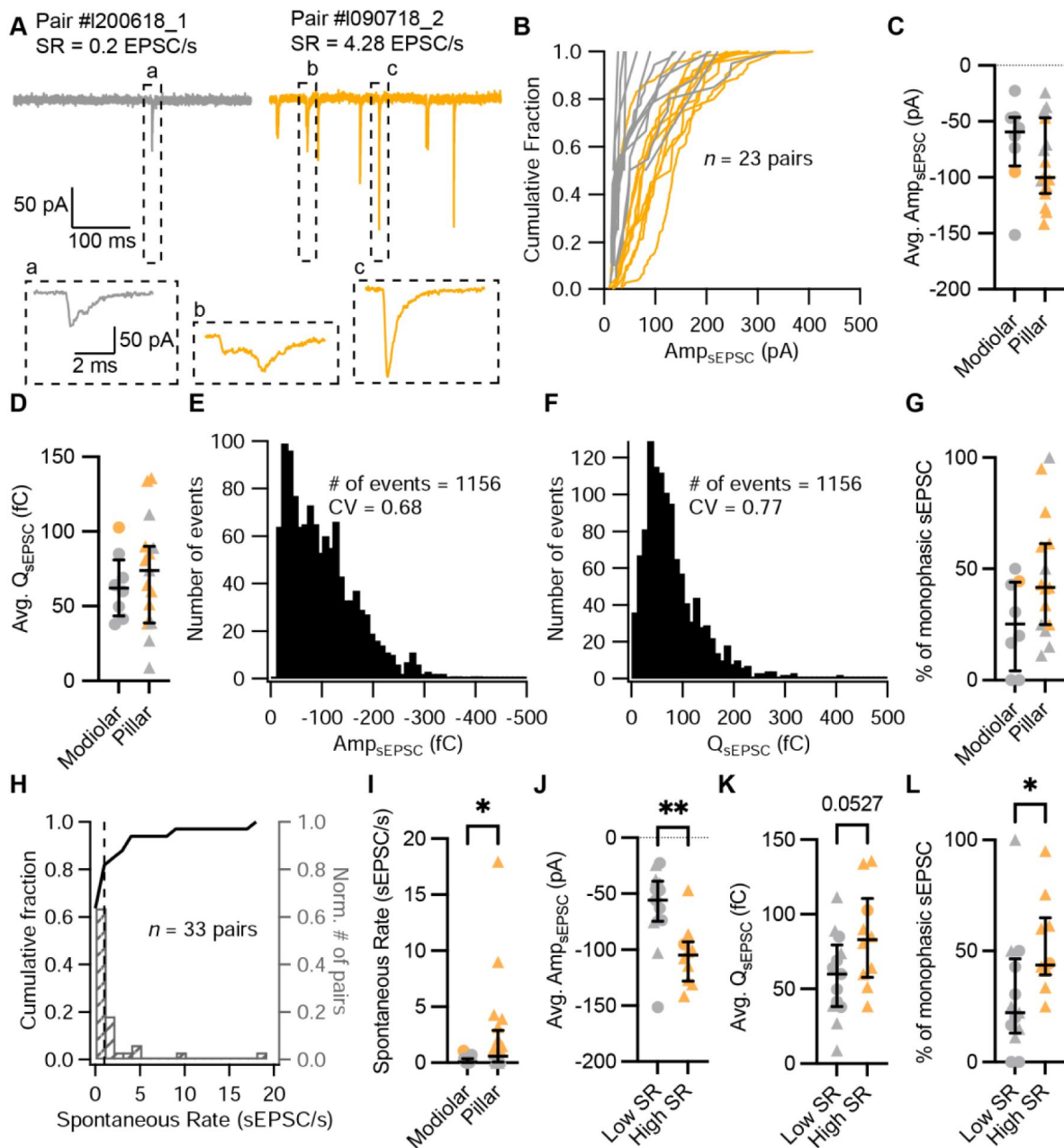


Figure 2.

Synapses with high spontaneous release have larger and more compact sEPSCs A.

Spontaneous EPSCs recorded in the absence of stimulation (i.e. IHC holding potential = -58 mV) from two exemplary paired recordings with different spontaneous rate (SR: grey for low SR, orange for high SR). 'Pair #' identifies individual paired recordings. Insets show the selected sEPSCs in an expanded time scale. (a,b) correspond to multiphasic sEPSCs, while (c) represents a typical monophasic sEPSC. **B.** Cumulative sEPSC amplitude plots for 23 paired synapses that had spontaneous release. **C-D.** Average sEPSC amplitude (C) and charge (D) from individual synapses recorded from the pillar or modiolar side of the IHC. **E-F.** Pooled sEPSC amplitude (E) and charge (F) distributions show a distinct peak at -40 pA and 40 pC, respectively. **G.** Percentage of monophasic sEPSCs in pillar and modiolar synapses. **H.** Cumulative fraction (left axis) and normalized histogram (right axis) of the spontaneous rate (bin size is 1 sEPSC/s) of 33 pairs. **I.** Pillar synapses had higher rates of sEPSCs ($p = 0.0311$, Mann-Whitney U test). **J-L.** High SR synapses had significantly larger sEPSC amplitudes (J; $p = 0.0042$, unpaired t-test), a tendency to bigger sEPSC charges (K; $p = 0.0527$, unpaired t-test) and higher percentages of monophasic sEPSCs ($p = 0.0185$, Mann-Whitney U test).

Whisker plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as Δ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.

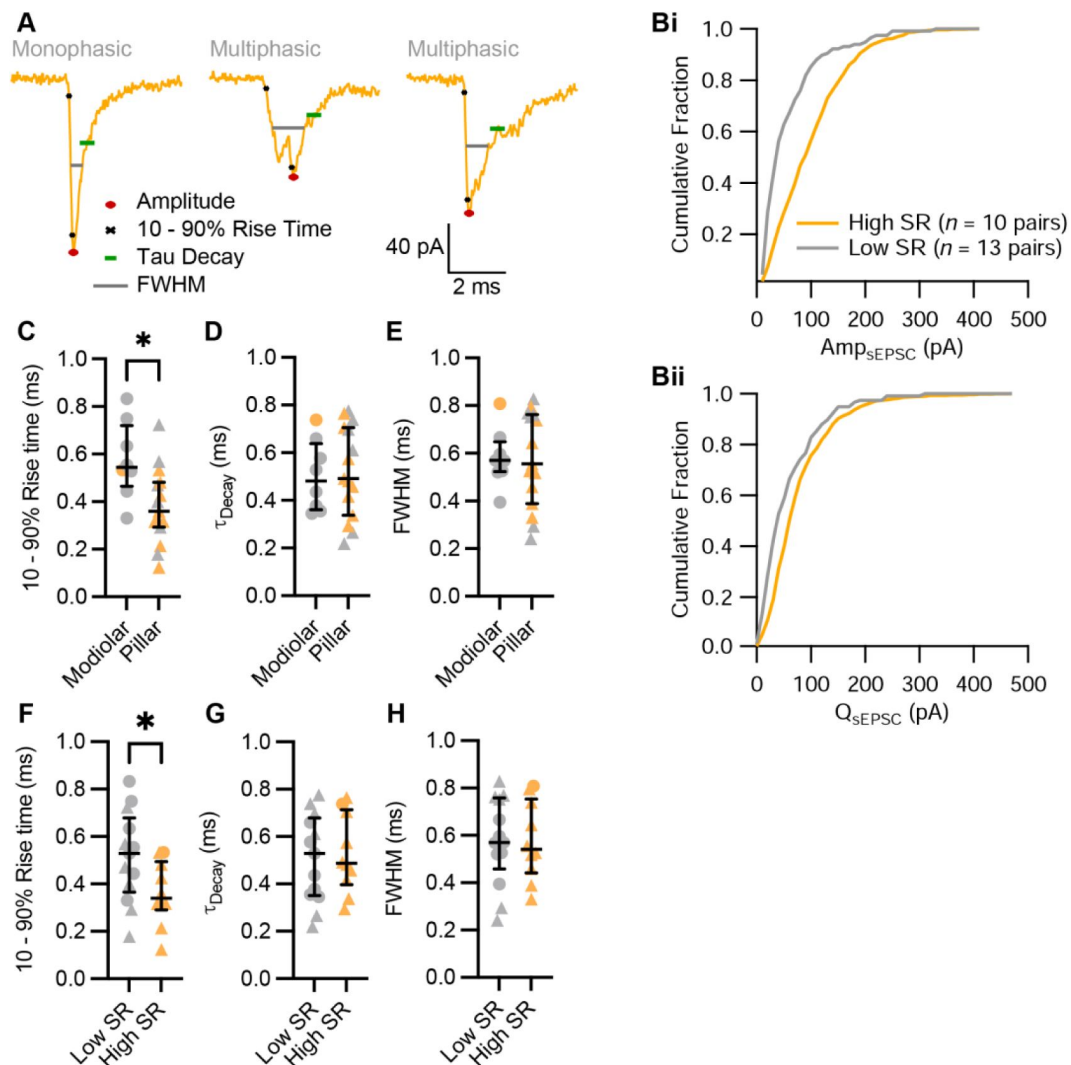


Figure 2-figure supplement 1.

Pillar and high SR synapses have sEPSCs with faster rising times.

A. Kinetics of sEPSCs, such as amplitude, 10-90% rise time, time constant of decay (τ_{decay}) and full-width half-maximum (FWHM), were calculated with Neuromatic (Rothman and Silver, 2018). **Bi-Bii.** Cumulative fraction of amplitude and charge of sEPSCs from low and high SR synapses. **C-E.** Pillar synapses had faster 10-90% rise times (C) than modiolar synapses ($p = 0.0111$, unpaired t-test), while their τ_{decay} (D) and FWHM (E) were comparable. **F-H.** High SR synapses had faster 10-90% rise times (F) than modiolar synapses ($p = 0.0420$, unpaired t-test), while their τ_{decay} (G) and FWHM (H) were comparable. Whisker plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as Δ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.

Evoked synaptic transmission differs between afferent synapses with high and low SR

Next, we compared the physiology of afferent synapses with high and low SR by adapting stimulation protocols routinely employed for *in vivo* characterization of sound encoding by SGNs. We used step depolarizations to emulate physiological receptor potentials given that mature IHCs of the “high frequency” mouse cochlea have graded receptor potentials that primarily represent the rectified envelope of an acoustic stimulus (i.e. the DC component) (Russell and Sellick, 1978 [\[1\]](#)).

i) Stimulus intensity encoding at IHC synapses

Sound intensity encoding by SGNs primarily relies on a spike rate code: the average discharge rate increases with the strength of the acoustic stimuli from threshold to saturation of the response. These so-called “rate level functions” are typically analyzed by fitting a sigmoidal function, of which the range of sound pressure level between 10 and 90% of the maximal discharge rate represents the operational or dynamic range (Sachs and Abbas, 1974 [\[2\]](#); Taberner and Liberman, 2005 [\[3\]](#); Winter et al., 1990 [\[4\]](#)). To understand stimulus intensity coding at mouse IHC synapses, we measured whole-cell IHC Ca^{2+} currents and the evoked EPSCs (eEPSCs) of SGNs in 31 paired recordings. We stimulated the IHC with 10 ms depolarizations to different potentials ranging from -70 to 0 mV in 5 mV steps (IV protocol; **Fig. 3A** [\[5\]](#)). We deemed it incompatible with a reasonable productivity of the technically challenging, low-throughput paired recordings to combine them with imaging of Ca^{2+} at single AZs. Therefore, this study relies on analysis of the presynaptic Ca^{2+} influx at the level of the whole IHC (i.e. summing over all synapses and a low density of extrasynaptic Ca^{2+} channels, (Frank et al., 2009 [\[6\]](#); Wong et al., 2014 [\[7\]](#)). IHCs with synapses classified as high ($n = 10$) or low SR ($n = 21$) had similar Ca^{2+} current-voltage (IV) relationships: comparable maximal Ca^{2+} currents (**Fig. 3Bi** [\[8\]](#); $p = 0.6939$, Mann-Whitney U test) elicited at similar potentials (**Fig. 3Bii** [\[9\]](#); $p = 0.1795$, unpaired t-test) and comparable reversal potentials (**Figure 3-figure supplement 1** [\[10\]](#), panel A; $p = 0.4034$, unpaired t-test). The fractional activation of Ca^{2+} channels was determined from the normalized chord conductance of the IHC. Fitting a Boltzmann function to these activation curves (**Fig. 3C** [\[11\]](#), upper panel), we obtained the voltages of half-maximal activation ($V_{\text{half Ca}}$) and the voltage-sensitivity of activation (slope) of the IHC Ca^{2+} channels.

“Release-stimulus intensity” curves, akin of an *ex vivo* representation of the SGN rate-level function, were constructed from the normalized Q_{EPSC} (**Figure 3-figure supplement 1** [\[12\]](#), panel C). The voltage dependence of synaptic vesicle release per active zone was approximated by the fit of a sigmoidal function to the individual release-intensity curves (**Fig. 3C** [\[13\]](#), lower panel). From these sigmoidal fits, we obtained voltage of 10%-maximal release ($Q_{10, \text{EPSC}}$), voltage of half-maximal release ($Q_{50, \text{EPSC}}$), voltage of 90%-maximal release ($Q_{90, \text{EPSC}}$) and the voltage sensitivity of release (slope). For two low SR paired recordings (**Fig. 3Dii** [\[14\]](#), grey dotted lines), a sigmoidal function did not properly fit the release-intensity curve (assessed by visual inspection) which led us to exclude them from the statistical analysis.

The whole-cell voltage dependence of Ca^{2+} influx was similar between IHCs contacted by high and low SR boutons: threshold of Ca^{2+} influx (**Fig. 3Ei** [\[15\]](#); $p = 0.2393$, unpaired t-test), $V_{\text{half Ca}}$ (**Fig 3Fi** [\[16\]](#); $p = 0.3479$, unpaired t-test) and voltage sensitivity of Ca^{2+} influx (**Figure 3-figure supplement 1** [\[17\]](#), panel B; $p = 0.3470$, unpaired t-test) did not differ significantly between IHCs contacted by low or high SR synapses. This seems to rule out a potential scenario in which the diverse SGN firing properties would be caused by varying average properties of Ca^{2+} channels among different presynaptic IHCs. $Q_{50, \text{EPSC}}$ (**Fig. 3Fii** [\[18\]](#)) of high SR synapses (-47.76 ± 1.4 mV; $n = 10$) was 6.76 ± 2.0 mV more negative compared to low SR synapses (-41.00 ± 1.2 mV; $n = 19$; $p = 0.0021$, unpaired t-test). Accordingly, high SR synapses had lower thresholds of release than low SR synapses (**Fig. 3Eii** [\[19\]](#), $Q_{10, \text{EPSC}}$ of -56.91 ± 1.5 mV (median -56.64 mV) vs. -47.39 ± 1.4 mV (median -48.89 mV); $p = 0.0001$, Mann-Whitney U test). The hyperpolarized shift was not significant for $Q_{90, \text{EPSC}}$ (**Figure 3-figure supplement 1** [\[20\]](#), panel D; $p = 0.1706$, unpaired t-test). The voltage sensitivity of release,

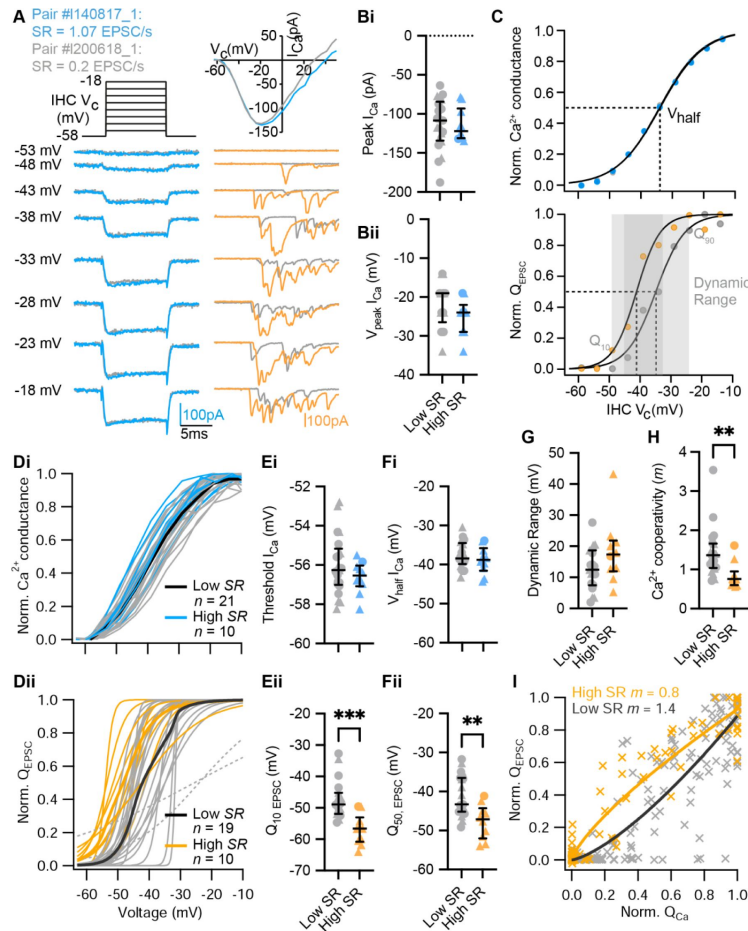


Figure 3.

High SR synapses have a more hyperpolarized voltage dependence and tighter Ca^{2+} channel coupling of synaptic release

A. IHC Ca^{2+} current (blue and gray) and eEPSCs (orange and gray) of a high and a low SR pair, respectively, in response to 10 ms depolarizations to different potentials ranging from -58 to -18 mV in 5 mV steps (upper left panel). The upper right panel shows the current-voltage relationships for the two pairs. **Bi-Bii.** The peak of whole-cell Ca^{2+} current (Bi) and the voltage eliciting maximum Ca^{2+} current (Bii) of IHCs were comparable between high and low SR synapses. **C.** Upper panel: Fractional activation of the Ca^{2+} channels (blue and gray data points from the examples shown in A) was obtained from the normalized chord conductance. Voltage of half-maximal activation ($V_{\text{half Ca}}$; dotted line) and sensitivity of the voltage dependence (slope k) were determined using a Boltzmann fit (black trace) to the activation curve. Lower panel: Release-intensity curve (orange and gray data points from the examples shown in A) was obtained from the Q_{EPSC} for each depolarization step. A sigmoidal function (black trace) was fitted to obtain the voltage of half-maximal synaptic release ($V_{\text{half QEPSC}}$; dotted line) and the voltage sensitivity of the release (slope), as well as the dynamic range for which the exocytosis changes from 10-90% (gray area). **Di-Dii.** Voltage dependence of whole-cell Ca^{2+} channel activation (activation curve; Di) and fits to release-intensity curves (Dii) for 31 synapses of low (< 1 sEPSC/s, gray) and high SR (≥ 1 sEPSC/s, orange). Averages (thick lines) and individual curves (thin lines) are overlaid. The release-intensity curve of two low SR pairs could not be fitted (grey dotted lines). **Ei-Fi.** The threshold of Ca^{2+} influx (Ei) and $V_{\text{half Ica}}$ (Fi) did not differ between low and high SR synapses. **Eii-Fii.** Voltage of 10% maximum release ($Q_{10 \text{ EPSC}}$; Eii; $p = 0.0001$, Mann-Whitney U test) and $V_{\text{half QEPSC}}$ (Fii; $p = 0.0021$, unpaired t-test) were significantly more hyperpolarized in high SR synapses. **G.** Dynamic range of release was comparable between low and high SR synapses. **H.** Ca^{2+} cooperativity (m) estimated from fitting a power function to the $Q_{\text{EPSC}} - Q_{\text{Ca}}$ relationship for each individual synapse was significantly lower in high SR synapses ($p = 0.0016$, Mann-Whitney U test). **I.** Scatter plot of normalized Q_{EPSC} versus the corresponding normalized Q_{Ca} . The solid lines is a least-squares fit of a power function ($Q_{\text{EPSC}} = a(Q_{\text{Ca}})^m$) to the data yielding $m_{\text{high SR}}$ of 0.8 (orange line) and $m_{\text{low SR}}$ of 1.4 (black line). Whisker plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as Δ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.

determined by a slope factor, tended to be higher in high SR (4.16 ± 0.75 mV) than in low SR (2.9 ± 0.35 mV) synapses without reaching statistical significance (**Fig. 3 – figure supplement 1** [↗](#), panel E; $p = 0.0940$, unpaired t-test;). The dynamic range, defined as the voltage range for which the exocytosis changes from 10-90% ($Q_{90\text{ EPSC}} - Q_{10\text{ EPSC}}$) tended to be larger for high SR synapses without reaching significance (**Fig. 3G** [↗](#); 18.30 ± 3.3 mV for high SR synapses vs 12.78 ± 1.5 mV for low SR synapses; $p = 0.0940$, unpaired t-test). The voltage dependence of Ca^{2+} influx and of synaptic vesicle release did not differ significantly when the synapses were grouped based on their topographical location at the IHC ($n = 12$ modiolar synapses vs $n = 17$ pillar synapses; **Figure 3–figure supplement 1** [↗](#), panels F-K). However, pillar SR synapses had a tendency to lower thresholds of release than modiolar synapses (**Figure 3–figure supplement 1** [↗](#), panel Gii; $Q_{10\text{ EPSC}}$ of -52.78 ± 1.8 mV (median -53.73 mV) vs. -47.68 ± 1.8 mV (median -49.30 mV); $p = 0.0725$, Mann-Whitney U test). Altogether, these results demonstrate that high SR synapses have a more hyperpolarized voltage-dependence of release.

Finally, we studied the apparent Ca^{2+} dependence of SV release during the aforementioned IV protocol, i.e., in the range of IHC receptor potentials. This protocol varies Ca^{2+} influx mainly via changing the channel open probability and to a lesser extent by changing the single channel current. We note that a supralinear intrinsic Ca^{2+} dependence of exocytosis in IHCs ($m \sim 3\text{--}4$) has been observed for IHCs of the cochlear apex in mice after hearing onset (Brandt, 2005 [↗](#); Özçete and Moser, 2021 [↗](#); Wong et al., 2014 [↗](#)). We related changes of release at individual synapses (ΔQ_{EPSC}) to the change of the integrated IHC Ca^{2+} influx (ΔQ_{Ca}). We fitted power functions ($Q_{\text{EPSC}} = a(Q_{\text{Ca}})^m$) to the relationships for individual synapses (**Figure 3–figure supplement 2** [↗](#)) and found Ca^{2+} cooperativities of $m < 2$ for all but 2 synapses. This result suggests a tight, Ca^{2+} nanodomain-like control of release sites by one or few Ca^{2+} channel(s). Interestingly, however, high SR synapses, on average, had significantly lower Ca^{2+} cooperativities than low SR synapses (**Fig. 3H** [↗](#); m_{highSR} of 0.8 ± 0.1 (median 0.75; $n = 10$) vs m_{lowSR} of 1.4 ± 0.1 (median 1.37; $n = 21$); $p = 0.0016$, Mann-Whitney U test). The fit to pooled normalized data of high and low SR synapses yielded the same Ca^{2+} cooperativities of m_{highSR} of 0.8 and m_{lowSR} of 1.4 (**Fig. 3I** [↗](#)). When grouped based on their modiolar or pillar location, pillar synapses showed significantly lower Ca^{2+} cooperativities than modiolar synapses (**Figure 3–figure supplement 1** [↗](#), panel L; m_{pillar} of 1.0 ± 0.08 (median 0.88; $n = 19$) vs m_{modiolar} of 1.6 ± 0.2 (median 1.3; $n = 12$); $p = 0.0202$, Mann-Whitney U test). Our findings indicate that most afferent IHC synapses of hearing mice employ a tight, Ca^{2+} nanodomain-like control of release sites by one or few Ca^{2+} channel(s) for physiological sound encoding. Yet, quantitative differences in coupling seem to exist between high SR/pillar synapses and low SR/modiolar synapses, whereby a control of SV release by ~ 1 Ca^{2+} channel prevails at high SR/pillar synapses.

ii) Synaptic vesicle pool dynamics at individual IHC active zones

In 13 of the 31 aforementioned paired recordings (6 classified as low SR and 7 as high SR; 2 belonging to modiolar and 11 to pillar synapses), we employed a forward masking paradigm to study synaptic vesicle (SV) pool dynamics of single afferent synapses. The forward masking paradigm (Harris and Dallos, 1979 [↗](#)) is commonly used for *in vivo* analysis of SGN spike rate adaptation and recovery from adaptation, which has been attributed to the depletion of readily releasable pool of SVs (RRP) and the recovery from depletion (Avissar et al., 2013 [↗](#); Frank et al., 2010 [↗](#); Furukawa and Matsuura, 1978 [↗](#); Goutman, 2017 [↗](#); Goutman and Glowatzki, 2007 [↗](#); Li et al., 2009 [↗](#); Moser and Beutner, 2000 [↗](#); Schroeder and Hall, 1974 [↗](#)). Typically, the *in vivo* protocol is applied at saturating sound pressure levels, which we aimed to mimic using strong step IHC depolarizations (to -18 mV from -58 mV) separated by different interstimulus intervals (ISI: 4, 16, 64 and 256 ms) (**Fig. 4A** [↗](#)). In analogy to the *in vivo* forward masking paradigm, the first stimulus - called *masker*, as it depresses the response to a subsequent stimulus when applied in rapid succession-had a duration of 100 ms. The second stimulus-denominated *probe*- lasted for 15 ms. The recordings included a time frame of 400 ms preceding the “masker” and 400 ms following the “probe”, and the interval between masker and masker was 20 s. Applied to recordings of eEPSCs,

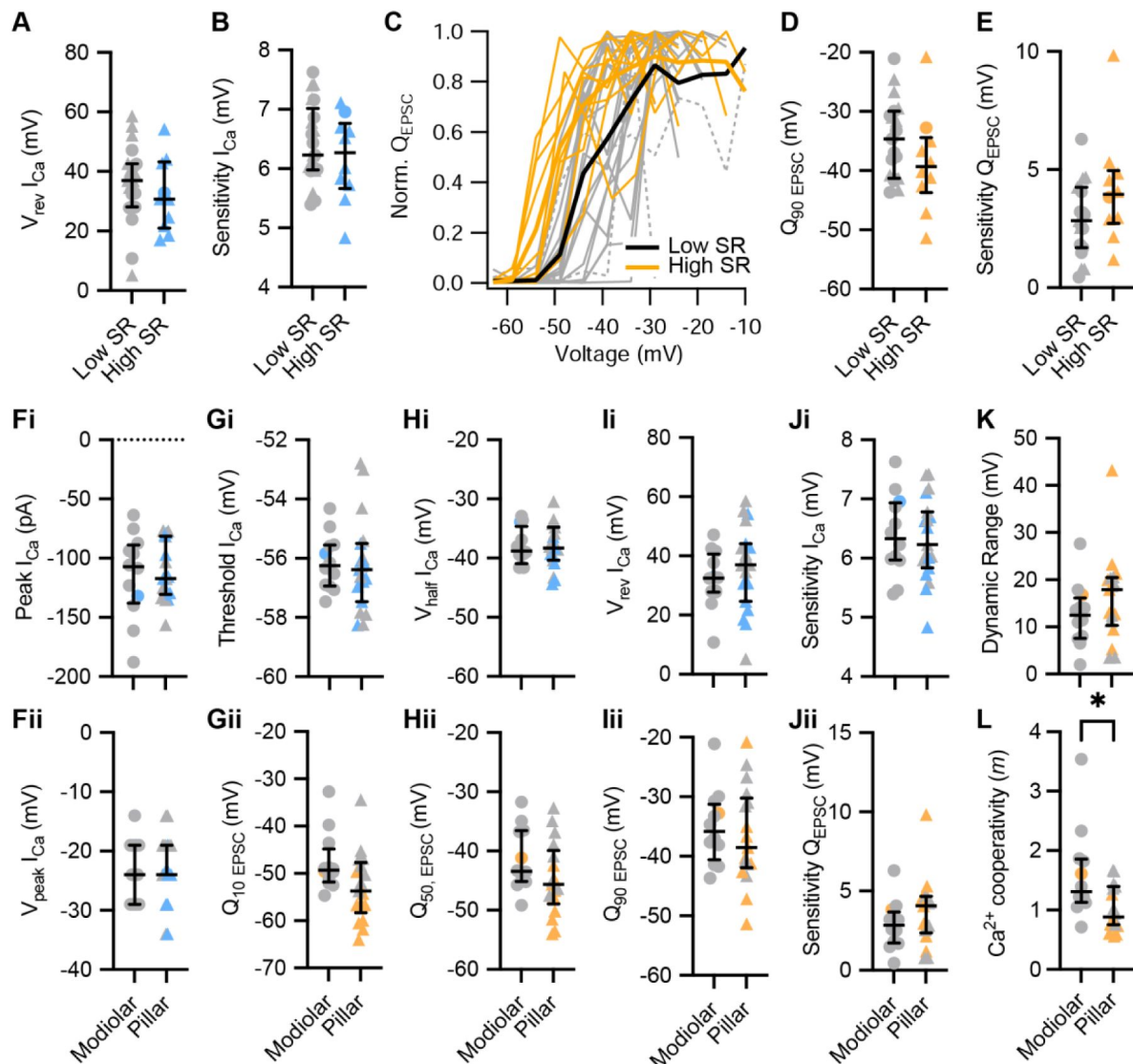


Figure 3 – figure supplement 1.

The voltage dependence of synaptic release does not differ significantly between modiolar and pillar synapses, but the Ca²⁺ dependence does

A, B. The reversal potential (A) and the voltage sensitivity of I_{Ca} (B) were comparable between low and high SR synapses. **C.** Triggered single active zone eEPSCs (release intensity curves) of 31 pairs of low and high SR. Averages (thick lines) and individual curves (thin lines) are overlaid. The release-intensity curve of two low SR pairs could not be fitted with the sigmoidal function (grey dotted lines). **D-E.** Q_{90} EPSC (D) and voltage sensitivity of the release (E) were comparable between high and low SR synapses. **Fi-Fii.** The peak of Ca²⁺ current (Fi) and the voltage eliciting maximum Ca²⁺ current (Fii) were comparable between modiolar and pillar synapses. **Gi-Ji.** The threshold of Ca²⁺ influx (Gi), V_{half} I_{Ca} (Hi), reversal potential of I_{Ca} (Ii) and voltage sensitivity of I_{Ca} (Ji) did not differ between modiolar and pillar synapses. **Gii-Jii.** Q_{10} EPSC (Gii), V_{half} Q_{EPSC} (Hii), Q_{90} EPSC (Iii) and voltage sensitivity of the release (Jii) were comparable between modiolar and pillar synapses. **K.** Dynamic range of release was comparable between modiolar and pillar synapse. **L.** Ca²⁺ cooperativity (m) estimated for each individual synapse was significantly lower in pillar synapses ($p = 0.0202$, Mann-Whitney U test). Whisker plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as Δ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.

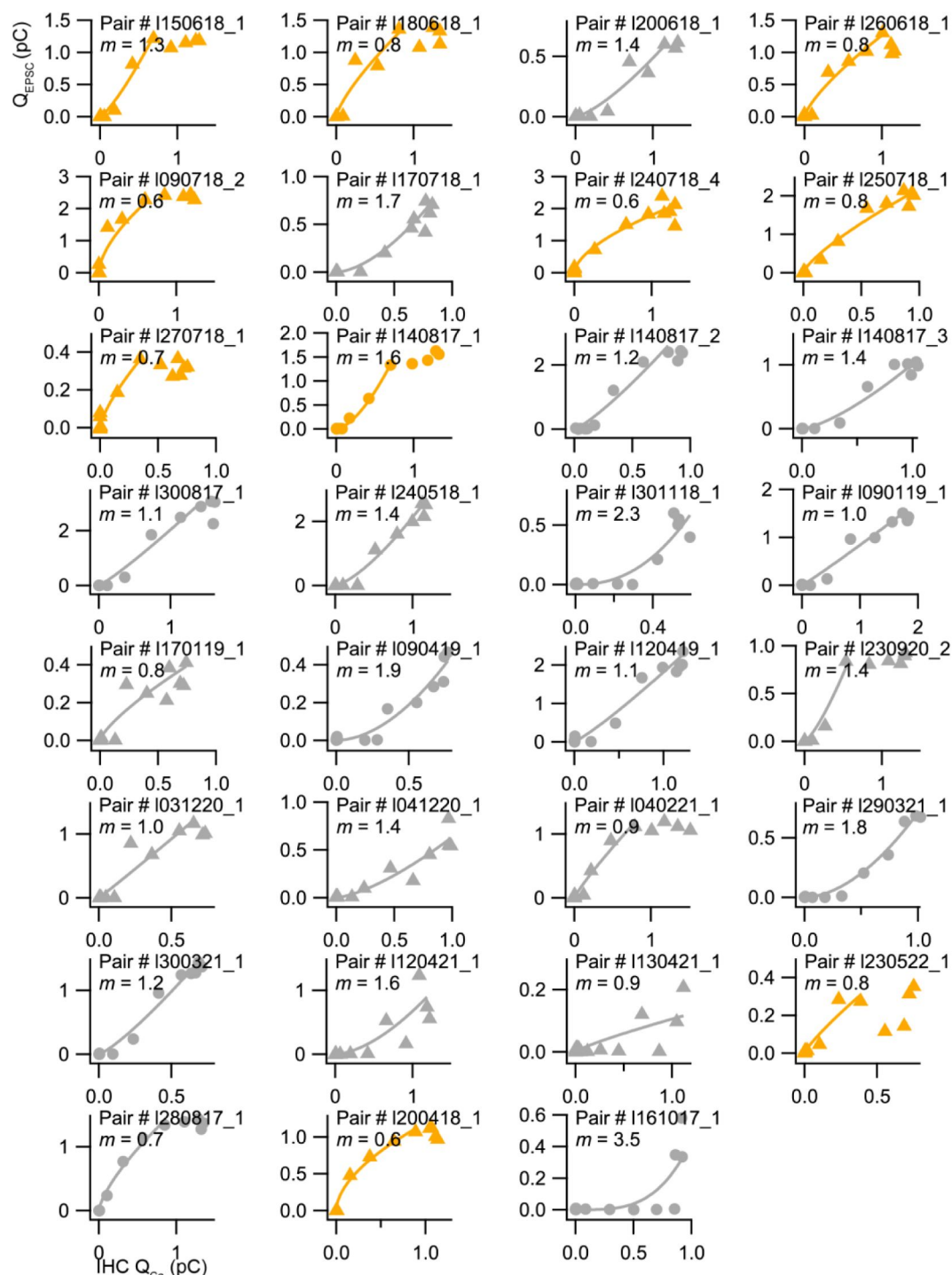


Figure 3 - figure supplement 2.

Apparent Ca^{2+} dependence of neurotransmitter release at individual synapses in the range of IHC receptor potentials

Scatter plots of the EPSC charges (Q_{EPSC}) vs. the corresponding Ca^{2+} current integrals (Q_{Ca}) for each individual synapse in response to 10 ms depolarizations from -58 to -18 mV. The solid line is a least-squares fit of a power function ($Q_{EPSC} = a(Q_{Ca})^m$) to each pair data.

the forward masking protocol provides experimental access to the initial RRP release rates, kinetics and extent of RRP depletion, sustained exocytosis, as well as recovery from RRP depletion. To accommodate the stochasticity of SV release from the RRP of IHC active zones, we run each protocol several times (≥ 3 to ≤ 20), which is routinely done for *in vivo* SGN physiology, but challenging *ex vivo* given the fragile and typically short-lived paired pre-and postsynaptic recordings (e.g. Goutman, 2017; Goutman and Glowatzki, 2007). Note that we did not employ cyclothiazide to inhibit AMPA receptor desensitization and reduce its contribution to postsynaptic eEPSC depression (Goutman, 2017), given the potential presynaptic effects of cyclothiazide in synaptic release (Diamond and Jahr, 1995; Dittman and Regehr, 1998).

For the analysis of evoked release dynamics, we focused on the response evoked by the masker. At the presynaptic level, there was no difference in the peak, initial and final IHC Ca^{2+} currents (I_{Ca} , **Figure 4-figure supplement 1**, panel A-C) and Ca^{2+} current charge (Q_{Ca} , **Fig. 4D**) between the recordings from high and low SR synapses. We calculated the synaptic delay from the onset of the masker to the onset of the eEPSC. High SR synapses had significantly shorter synaptic delays (**Fig. 4B**), with mean latencies of the first eEPSC after stimulus onset of 1.21 ± 0.09 ms (median 1.18) compared to 3.34 ± 1.03 ms (median 2.09) in low SR pairs ($p = 0.0221$, Mann-Whitney U test). This result was corroborated in a bigger sample size, when we compared the synaptic latencies of the 31 pairs to 10 ms pulses to -18 mV (**Figure 4-figure supplement 1**, panel D; latency of 1.19 ± 0.14 ms (median 1.14) in high SR compared to 2.57 ± 0.48 ms (median 1.79) in low SR synapses; $p = 0.0101$, Mann-Whitney U test). The latency jitter (measured as the standard deviation of the eEPSC latencies; **Fig. 4C**) was also significantly smaller in high SR synapses compared to low SR synapses (0.38 ± 0.07 ms (median 0.41) vs 1.67 ± 0.72 ms (median 0.98), respectively; $p = 0.0012$, Mann-Whitney U test). Additionally, high SR synapses had significantly larger peak amplitudes of the masker-evoked EPSCs which reports the initial release from the RRP (**Figure 4-figure supplement 1**, panel E; -421.4 ± 54.22 pA for high SR synapses vs -240.2 ± 23.25 pA for low SR synapses; $p = 0.0147$, unpaired t-test). In contrast, the charge of masker-evoked EPSCs ($Q_{\text{EPSC MASKER}}$) was more comparable (**Figure 4-figure supplement 1**, panel F; 4.818 ± 0.88 pC for high SR vs 3.223 ± 0.47 pC for low SR pairs; $p = 0.1574$, unpaired t-test). We fitted the first 50 ms of the $Q_{\text{EPSC MASKER}}$ with the sum of a single exponential and a line function in order to analyze RRP depletion and sustained release (**Figure 4-figure supplement 1**, panel G; **Fig. 4G** and dashed lines in **Fig. 4D**). The amplitude of the exponential component (A_1), thought to reflect RRP exocytosis, was not different between high and low SR synapses (**Figure 4-figure supplement 1**, panel H; $p = 0.4092$, unpaired t-test). Likewise, the slope of the linear component, reflecting sustained exocytosis, did not differ significantly between the two groups (**Figure 4-figure supplement 1**, panel I; $p = 0.1807$, Mann-Whitney U test).

To quantify synaptic release in terms of SVs, we divided $Q_{\text{EPSC MASKER}}$ by the mean Q_{sEPSC} recorded for each pair. This builds on our assumption that each sEPSC corresponds to a unitary release event (“univesicular mode of release” (Chapachnikov et al., 2014; Grabner and Moser, 2018; Huang and Moser, 2018)). The quantal content (RRP size in #SV) was comparable between high and low SR synapses (**Fig. 4F**, 13.99 ± 2.239 SVs vs 14.28 ± 4.839 SVs, respectively; $p = 0.9553$, unpaired t-test). However, the RRP depleted significantly faster in high SR synapses (**Fig. 4G**): $\tau_{\text{RRP depletion}}$ was 6.347 ± 1.096 ms (median 6.13) for high SR vs. 20.88 ± 5.063 ms (median 19.93) for low SR synapses ($p = 0.0140$, Mann-Whitney U test). Accordingly, high SR synapses showed significantly higher initial release rates compared to low SR synapses (**Fig. 4H**; 2651 ± 660.0 SV/s vs. 927.4 ± 307.4 SV/s; $p = 0.0472$, unpaired t-test), which given the comparable RRP size, indicates a higher release probability of the high SR rare synapses. Moreover, high SR synapses showed stronger depression of the release rate (**Fig. 4J**; the ratio of initial/sustained release rate was 5.941 ± 0.916 for high SR pairs and 2.023 ± 0.6251 for low SR pairs; $p = 0.006$, unpaired t-test), despite comparable sustained release rates (**Fig. 4I**; $p = 0.9258$, unpaired t-test).

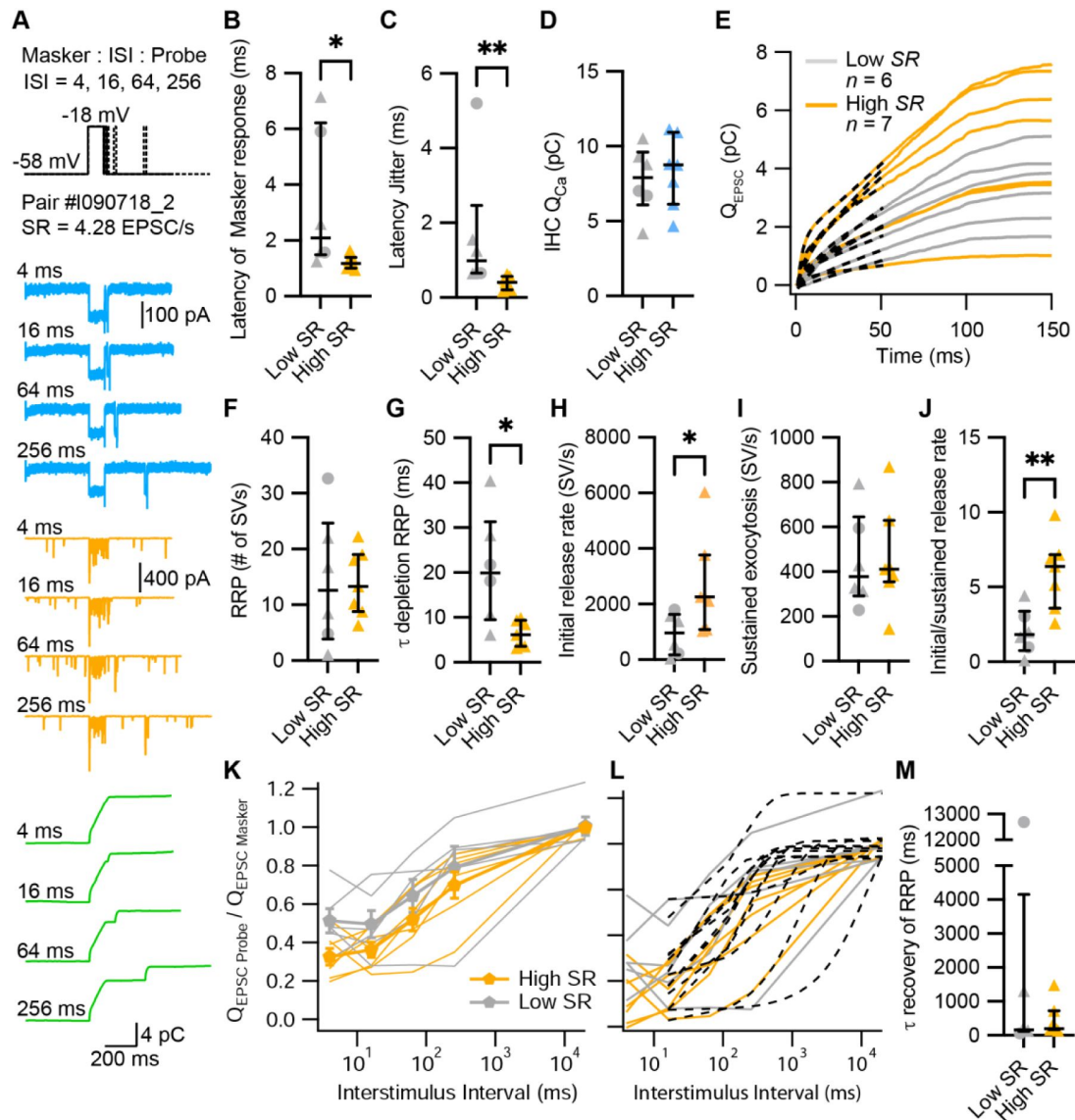


Figure 4.

High SR synapses have shorter synaptic delay and higher initial release rates

A. Representative Ca^{2+} currents (blue), eEPSCs (light orange) and eEPSC charges (Q_{EPSC} , green) to the “forward masking” protocols used to study depletion and recovery of RRP. The stimulus (top panel) consists of two sequential voltage steps (“masker” and “probe”) separated by different interstimulus intervals (ISI in ms). **B.** Latencies of the evoked EPSCs ($\text{EPSC}_{\text{onset}} - \text{Masker}_{\text{onset}}$) were significantly shorter in high SR than low SR synapses ($p = 0.0221$, Mann-Whitney U test). **C.** High SR synapses also had less latency jitter ($p = 0.0012$, Mann-Whitney U test). **D.** Presynaptic IHC Ca^{2+} charge (Q_{Ca}) during the masker stimuli was comparable regardless the spontaneous rate of the recorded postsynaptic bouton. **E.** Pool depletion dynamics were studied by fitting the sum of a single exponential and a line function (black discontinuous line) to the first 50 ms of the average Q_{EPSC} trace in response to the masker stimulus. **F–J.** RRP, time constant (τ) of depletion, initial release rate and sustained release were calculated from the fits and the mean Q_{EPSC} for each pair. High SR synapses depleted the RRP with faster time constants (**G**; $p = 0.0140$, Mann-Whitney U test) and reached higher initial release rates (**H**; $p = 0.0472$, unpaired t-test) followed by a stronger adaptation (**J**; $p = 0.0058$, unpaired t-test). **K.** Recovery from RRP depletion shown as ratio of $Q_{\text{EPSC probe}}$ and $Q_{\text{EPSC masker}}$ (mean $\pm$ sem) during the first 10 ms of the stimulus. **L.** Single exponential fits from 16 ms to 20000 ms (black dotted lines) to estimate the recovery kinetics from RRP depletion. **M.** Time constant of recovery from RRP depletion obtained from fits in **L**.

Whisker plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as $\triangle$ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.

Finally, we determined RRP recovery from depletion using the ratio $Q_{\text{EPSC Probe}}/Q_{\text{EPSC Masker}}$, whereby Q_{EPSC} for masker and probe was estimated for the first 10 ms of stimulation. **Figure 4K** plots the ratio for each interstimulus interval (ISI), including the masker-to-masker interval. Surprisingly, and contrary to a previous report in auditory bullfrog synapses (Cho et al., 2011), approximately half of the synapses showed a lower ratio at 16 ms than at 4 ms regardless of their SR. Therefore, to determine the kinetics of recovery from RRP depletion, we fitted a single exponential to the recovery data over the ISI range from 16 ms to 20 s (**Fig. 4L**). We did not observe a significant difference between high SR and low SR synapses (**Fig. 4M**; $\tau_{\text{recovery RRP}}$: 451.5 ± 187.0 ms (median 192.7) vs 2416 ± 2064 ms (median 157.9), respectively; $p = 0.5338$, Mann-Whitney U test). In 4 high SR and 4 low SR synapses, spontaneous activity was resumed shortly after the offset of the probe. The time to this first sEPSC took longer in high SR synapses compared to low ones (**Figure 4-figure supplement 1**, panel J; 185.2 ± 25.25 ms (median 161.6) vs 104.5 ± 24.10 ms (median 108.4), respectively; $p = 0.0286$, Mann-Whitney U test), again indicating stronger synaptic depression of high SR synapses.

Discussion

Much of the information on synaptic sound encoding at afferent IHC-SGN synapses has been obtained from either juxtacellular recordings of SGN firing *in vivo* or from *ex vivo* patch-clamp recordings. Yet, it has remained difficult to reconcile those *in vivo* and *ex vivo* results and to establish a unified account of sound intensity coding in the auditory nerve given differences in experimental conditions, animal models and protocols employed. Here, we biophysically characterized the heterogeneous function of afferent SGN synapses in hearing mice with reference to their rate of spontaneous transmission (SR) as a surrogate of SGN SR that informs their functional properties. We performed paired pre- and post-synaptic patch-clamp recordings from single IHC synapses of hearing mice under near-physiological conditions using protocols adapted from *in vivo* characterization of SGN's response properties. Using this approach, we were able to distinguish synapses with low and high SR, which we propose to provide the input into low and high SR SGNs. We found that about 90% of high SR synapses were located at the pillar side of the IHC. High SR synapses had larger sEPSCs with a more compact (monophasic) waveform, lower voltage-thresholds of release, shorter synaptic delays, tighter coupling of release sites to Ca^{2+} channels as well as higher initial release rates and shorter RRP depletion time constants. We conclude that high SR synapses exhibit higher release probability which likely reflects the tighter coupling of Ca^{2+} channels and release sites. RRP size, rate of sustained exocytosis and kinetics of RRP recovery from depletion were comparable between high and low SR synapses.

Diversity of spontaneous release and their topographical segregation

The SR range observed in our paired recordings from mouse afferent synapses (0 to 18 EPSC/s) agrees with previous reports of bouton recordings from rat synapses (p15-p17, Wu et al., 2016). However, the maximum rate is considerably smaller than those recorded *in vivo* from single SGNs [up to 60 spikes/s in p14-p21 mice (Wong et al., 2013) and up to 120 spikes/s in 8-17 weeks old mice (Taberner and Liberman, 2005)]. Given that less than 35% of murine SGNs have *in vivo* SRs above 20 spikes/s (Taberner and Liberman, 2005), our lower maximum rate could also originate from a sampling bias. In addition, the larger, faster and more compact sEPSC observed in high SR synapses would increase the speed and timing of the generated spike(s) (Rutherford et al., 2012), perhaps allowing for several spikes to be generated and leading to a further increase in the SGN spike SR. Other important factors that might explain the higher SR reported *in vivo* include: i) the intrinsic biophysical properties of SGNs which could further expand the firing rate distribution (Markowitz and Kalluri, 2020); ii) the developmental recruitment of high SR SGNs with age (Niwa et al., 2021; Romand, 1984; Walsh and McGee, 1987; Wong et al., 2013; Wu et al., 2016);

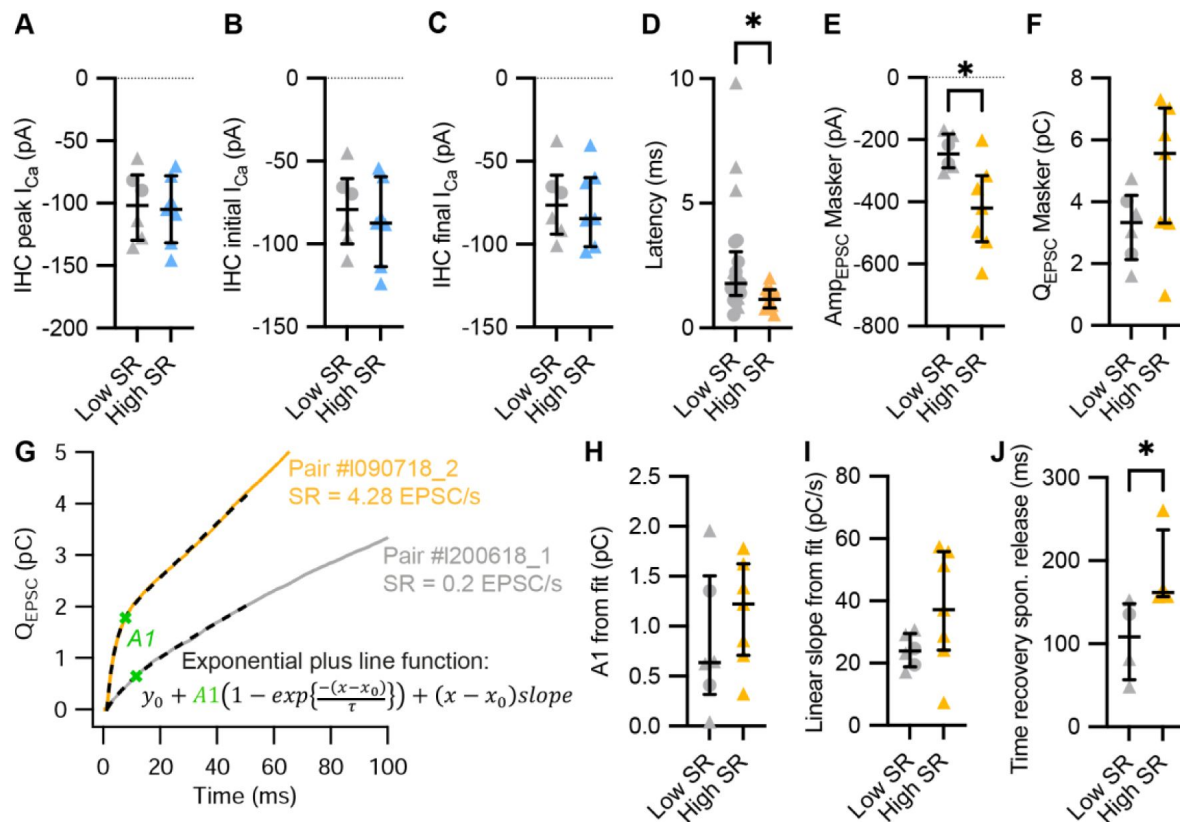


Figure 4 – figure supplement 1.

Parameters of synaptic vesicle pool dynamics in forward masking protocols

A–C. Presynaptic peak (A), initial (B) and final (C) Ca^{2+} current (I_{Ca}) during the masker stimuli were comparable regardless the SR of postsynaptic bouton. **D.** Latencies of the evoked EPSCs were also significantly shorter in high SR than low SR synapses when compared in a bigger sample size (31 pairs from Fig. 4; $p = 0.0101$, Mann-Whitney U test). **E–F.** High SR synapses had significantly larger amplitudes of evoked EPSCs in response to the masker stimulus (E; $p = 0.0147$, unpaired t-test), while the EPSC charge (Q_{EPSC}) did not differ (F). **G.** Exemplary average Q_{EPSC} response of a high SR (orange) and a low SR (gray) synapse to the masker stimuli. We fitted an exponential plus line function to the first 50 ms of the response (discontinuous lines) to study SV pool depletion dynamics. From these fits, we can retrieve information about RRP size (amplitude of the exponential component, A1), RRP depletion time constant (τ of the exponential component) and sustained release (linear slope of the line component). **H–I.** Amplitude (H) and linear slope (I) obtained from the fits of the exponential plus line function to the Q_{EPSC} from individual pairs (Fig. 4E). **J.** The spontaneous release after the probe offset recovered slower in high SR synapses.

Whisker plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as Δ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR $\geq$ 1 sEPSC/s.

and iii) compromised physiological conditions as well as a deviation of the chosen IHC holding potential from the IHC resting potential despite our efforts to mimic physiological conditions during the *ex vivo* recordings.

In analogy to the pioneering finding of a synaptic segregation of cat SGNs according to SR along the pillar–modiolar axis of IHCs (Lieberman, 1982 [↗](#); Merchan-Perez and Liberman, 1996 [↗](#)), we found that about 90% of high SR were located on the pillar IHC side. Yet, not all the synapses of the pillar IHC side had high SR, which agrees with a recent study of molecularly tagged SGNs (Siebald et al., 2023 [↗](#)). These findings suggest that high frequency and large amplitude of sEPSCs occur predominantly in synapses with smaller ribbons and AZs, opposing results from retinal cells in which smaller ribbons resulted in reduced frequency and amplitude of EPSCs (Mehta et al., 2013 [↗](#)). It is important to point out that our modiolar/pillar classification is less precise than that of other studies in which the synapse position was quantitatively assigned (Frank et al., 2009 [↗](#); Liberman et al., 2011 [↗](#); Ohn et al., 2016 [↗](#); Özçete and Moser, 2021 [↗](#)). Moreover, other studies also support an overall pillar-modiolar gradient with “salt and pepper” intermingling of synaptic properties rather than their strict segregation (Ohn et al., 2016 [↗](#); Özçete and Moser, 2021 [↗](#)).

Similar to previous reports (e.g. Chapochnikov et al., 2014 [↗](#); e.g. Glowatzki and Fuchs, 2002 [↗](#); Grant et al., 2010 [↗](#); Huang and Moser, 2018 [↗](#); Rutherford et al., 2012 [↗](#)), we found a high variability in the waveform and amplitude of sEPSCs between synapses, which apparently do not strictly depend on the topographical location of the synapse (the present work and (Niwa et al., 2021 [↗](#))). Yet, high SR synapses had larger and more compact (monophasic) sEPSCs. The difference in the percentage of monophasic sEPSCs in low and high SR synapses could arise from variability in the fusion pore dynamics on the way to vesicle fusion and/or on the number of SVs released in timely manner. In the view of the multivesicular hypothesis of spontaneous release (Glowatzki and Fuchs, 2002 [↗](#); Li et al., 2009 [↗](#); Niwa et al., 2021 [↗](#); Schnee et al., 2013 [↗](#)), SVs of a modiolar AZ might fuse in an uncoordinated manner, creating a EPSCs with a less compact waveform. Alternatively and our favorite hypothesis, each sEPSC corresponds to a unitary release event (“univesicular mode of release” (Chapochnikov et al., 2014 [↗](#); Grabner and Moser, 2018 [↗](#); Huang and Moser, 2018 [↗](#); Young et al., 2021 [↗](#))) that has the capacity to drive action potential firing (Rutherford et al., 2012 [↗](#)). In the framework of the univesicular hypothesis of spontaneous release, the flickering of the fusion pore, prior to or instead of full collapse fusion of the SV, might be favored in low SR synapses, leading to a lower percentage of monophasic sEPSCs. Such heterogeneity of fusion pore dynamics has been reported in chromaffin cells, calyx of Held and hippocampal neurons (Chang et al., 2021 [↗](#); Henkel et al., 2019 [↗](#); Shin et al., 2020 [↗](#), 2018 [↗](#)).

Candidate mechanisms distinguishing evoked release at low and high SR synapses

The temporal and quantal resolution offered by paired recordings allowed us to analyze the biophysical properties of evoked synaptic transmission in relation to the SR of the given synapse. In an intriguing resemblance with *in vivo* evoked firing properties of high SR SGNs (Bourien et al., 2014 [↗](#); Buran et al., 2010 [↗](#); Relkin and Doucet, 1991 [↗](#); Rhode and Smith, 1985 [↗](#); Taberner and Liberman, 2005 [↗](#)), high SR synapses showed lower voltage (–sound pressure *in vivo*) thresholds of synaptic transmission (–firing *in vivo*), shorter and less variable synaptic latencies (–first spike latencies *in vivo*), and higher initial release rates (–onset firing rate *in vivo*). In addition, we found stronger synaptic depression at high SR synapses, which agrees well with the finding of a greater ratio of peak to adapted firing rate in high SR SGNs recorded *in vivo* (Taberner and Liberman, 2005 [↗](#)). These results support the hypothesis that IHC synaptic heterogeneity (Frank et al., 2009 [↗](#); Hua et al., 2021 [↗](#); Ohn et al., 2016 [↗](#); Özçete and Moser, 2021 [↗](#); Reijntjes et al., 2020 [↗](#)) contributes to the diversity of spontaneous and sound-evoked SGN firing.

How do high *SR* synapses with likely smaller ribbons and lower maximal Ca^{2+} influx achieve a shorter latency and higher initial release rate? Our hypothesis is that, in high *SR* synapses, a more hyperpolarized Ca^{2+} channel activation (Ohn et al., 2016) in combination with tighter coupling between the Ca^{2+} channels and the Ca^{2+} sensor of fusion (this work and (Özçete and Moser, 2021)) would enable a faster response with a greater initial SV release probability for a given stimulus. Interestingly, genetic disruptions that shift the voltage-dependence have a greater impact on the *in vivo* distribution of *SR* than mutations that change the maximal synaptic Ca^{2+} influx (Jean et al., 2018; Ohn et al., 2016). The spatial coupling of the Ca^{2+} channel to the SV release site has also been shown to greatly affect release probability in other synapses (Eggermann et al., 2012; Fekete et al., 2019; Moser et al., 2019; Rebola et al., 2019). Other factors that affect release probability include variations in the number of open Ca^{2+} channels at the AZ (Gratz et al., 2019; Holderith et al., 2012; Scimemi and Diamond, 2012; Sheng et al., 2012; Wong et al., 2013) and the fusion competence of the SV (Klenchin and Martin, 2000), including the priming and docking state (Lin et al., 2022; Neher and Brose, 2018). The ensuing Ca^{2+} influx in pillar synapses might facilitate Ca^{2+} channels and priming of SVs (Cho and von Gersdorff, 2012; Goutman and Glowatzki, 2011, 2007; Michalski et al., 2017; Moser and Beutner, 2000; Pangrsic et al., 2015; Schnee et al., 2011; Spassova et al., 2004) and contribute to the observed results in high *SR* synapses. Regarding the fusion competence of SVs, it is unknown whether modiolar and pillar synapses exhibit different numbers of docked and primed SVs. To date, ultrastructural studies that resolve docked and tethered SVs have not address the topographical location of the AZ in the murine IHC (Chakrabarti et al., 2022, 2018).

Besides SV release probability, RRP size co-determines neurotransmitter release. Our estimated RRP of about 14 SVs in both high and low *SR* synapses compares well to prior estimates obtained using *ex vivo* electrophysiology (10-40 SVs: Refs. (Goutman and Glowatzki, 2007; Jean et al., 2018; Johnson et al., 2005; Khimich et al., 2005; Moser and Beutner, 2000; Pangrsic et al., 2010; Schnee et al., 2005), model-based analysis of SGN firing (4-40 SVs (Frank et al., 2010; Jean et al., 2018; Peterson et al., 2014)) and electron microscopy (10-16 SVs within 50 nm of the presynaptic membrane (Chakrabarti et al., 2018; Frank et al., 2010; Graydon et al., 2011; Kantardzhieva et al., 2013; Khimich et al., 2005; Pangrsic et al., 2010)). However, previous reports based on electron microscopy (Kantardzhieva et al., 2013; Merchan-Perez and Liberman, 1996; Michanski et al., 2019) suggested larger pools of SVs at modiolar synapses, while our electrophysiological estimate of RRP size was comparable between low and high *SR* synapses. This finding argues against a strong contribution of RRP size to the observed differences in neurotransmitter release. However, higher release probability with comparable RRP size explains higher initial release rates, which likely explain the faster and temporally more precise postsynaptic depolarization that is likely to turn into shorter first spike latencies and lower first spike latency jitter (this study and (Buran 2010).

Finally, the heterogeneity in the functional properties of IHC synapses could also arise from molecular heterogeneity of the AZ. In central glutamatergic synapses, molecular heterogeneity of synaptic proteins plays a critical role in the modulation of SV release probability and priming state (Neher and Brose, 2018; Wichmann and Kuner, 2022). For instance, differential isoforms of priming factors and scaffold proteins have been suggested to tune the functional synaptic diversity of central synapses (Fulterer et al., 2018; Rebola et al., 2019; Rosenmund et al., 2002). Cochlear IHCs have an unconventional fusion machinery that appears to work without neuronal SNAREs (Nouvian et al., 2011) (but see Calvet et al., 2022) and priming factors such as Munc13 and CAPS (Vogl et al., 2015). Therefore, future studies will need to determine the molecular nanoanatomy underlying the specific AZ nanophysiology and functional synaptic heterogeneity at IHCs. Promising candidates include RBPs (Butola et al., 2021; Grauel et al., 2016; Krinner et al., 2021, 2017; Petzoldt et al., 2020), RIMs (Jung et al., 2015; Maria M. Picher et al., 2017), and Septin (Fekete et al., 2019; Yang et al., 2010).

Challenges for relating synaptic and neural response properties

Next to providing support for the presynaptic hypothesis of functional SGN diversity, the present study also highlights some of the challenges met when aiming to bridge the gap between presynaptic hair cell function and neural sound encoding. Despite major efforts undertaken to match experimental conditions and protocols, it remains difficult to reconcile some findings of *ex vivo* and *in vivo* physiology. Parameters such as RRP size (~# spikes of the rapidly adapting component of firing), sustained exocytosis (~adapted firing rate *in vivo*), recovery of spontaneous and evoked release (~recovery from forward masking *in vivo*) did not differ among our high and low SR synapses, and contrasts with *in vivo* data (e.g. Refs. (Bourien et al., 2014 [DOI](#); Buran et al., 2010 [DOI](#); Relkin and Doucet, 1991 [DOI](#); Rhode and Smith, 1985 [DOI](#); Taberner and Liberman, 2005 [DOI](#))).

Of particular interest is that the dynamic ranges and slope of release-intensity relationship of high and low SR synapses diverge from the expectations if assuming that high SR SGNs are driven by high SR synapses. High SR synapses tended to show broader dynamic ranges with shallower slopes, while, *in vivo*, high SR SGNs show smaller dynamic ranges and steeper slopes than the low SR ones (Ohlemiller et al., 1991 [DOI](#); Winter et al., 1990 [DOI](#)). Could this reflect the non-linear saturating properties of the basilar membrane (Sachs et al., 1989 [DOI](#); Sachs and Abbas, 1974 [DOI](#); Yates et al., 1990 [DOI](#)) (discussed in Ref. (Ohlemiller et al., 1991 [DOI](#))) which might widen the rate level function of low SR SGNs? Or is it due to a partial depletion of the “standing” RRP (i.e. the occupancy of the RRP release sites with a fusion-competent SV (Moser, 2020 [DOI](#); Pangrsic et al., 2010 [DOI](#))) at high SR synapses *in vivo*? It remains to be determined whether this and the other aforementioned differences between our data and *in vivo* reports could be attributed to mechanisms downstream of glutamate release and AMPA receptor activation. The possible mechanisms include but are not limited to: i) different spike rates due to diverse EPSC waveforms (Rutherford et al., 2012 [DOI](#)); ii) differences in SGN excitability (Crozier and Davis, 2014 [DOI](#); Markowitz and Kalluri, 2020 [DOI](#); Smith et al., 2015 [DOI](#)) due to heterogeneous molecular (Petitpré et al., 2020 [DOI](#), 2018 [DOI](#); Shrestha et al., 2018 [DOI](#); Sun et al., 2018 [DOI](#)) and morphological profiles (Liberman, 1980 [DOI](#); Merchan-Perez and Liberman, 1996 [DOI](#); Tsuji and Liberman, 1997 [DOI](#)); and iii) differences in efferent innervation of SGNs (Hua et al., 2021 [DOI](#); Liberman, 1990 [DOI](#); Ruel et al., 2001 [DOI](#); Wu et al., 2020 [DOI](#); Yin et al., 2014 [DOI](#)). Certainly, caution is to be applied for the comparison of *ex vivo* and *in vivo* data due to partial disruption of the physiological milieu despite our efforts to maintain near-physiological conditions. This also extends to the alteration of efferent signaling in our *ex vivo* preparation.

Clearly more work is needed to elucidate the mechanisms of SGN firing diversity in the cochlea. Ideally, future studies will combine *in vivo* and *ex vivo* experiments, such as combining physiological SGN characterization with neural backtracing and synaptic morphology of labelled SGNs using volume imaging of afferent and efferent connectivity (Hua et al., 2021 [DOI](#)). Moreover, combining optogenetic IHC stimulation with imaging of SGN activity could provide higher throughput and serve posthoc morphology. Finally, paired patch-clamp recordings, as done in the present study, could be combined with SGN subtype-specific molecular labeling, fiber tracing and immunolabeling to further relate synaptic transmission and SGN neurophysiology.

Materials and Methods

Animals and tissue preparation

c57BL/6N mice of either sex between postnatal day 14-23 (p14-23) were used. The animal handling and experiments complied with national animal care guidelines and were approved by the University of Göttingen Board for animal welfare and the Animal Welfare Office of the State of Lower Saxony. Animals were sacrificed by decapitation and the cochleae were extracted in modified Hepes Hank's solution containing: 5.36 mM KCl, 141.7 mM NaCl, 1 mM MgCl₂·6H₂O, 0.5 mM MgSO₄·7H₂O, 10 mM HEPES, 0.5 mg/ml L-glutamine, and 1 mg/ml D-glucose (pH 7.2,

osmolarity of ~300 mOsm). The apical coil of the organ of Corti was dissected and placed under a grid in the recording chamber. Pillar or modiolar supporting cells were removed using soda glass pipettes in order to gain access to the basolateral face of the IHCs and to the postsynaptic boutons of type I SGNs. Dissection of the organ of Corti and cleaning of the supporting cells were performed at room temperature (20–25°C).

Electrophysiological recordings

Pre- and postsynaptic paired patch clamp recordings were performed at near physiological temperature (32–37°C) using an EPC-9 amplifier (HEKA electronics) (Fig. 1). Patch electrodes were positioned using a PatchStar micromanipulator (Scientifica, UK). Whole-cell recordings from IHCs were achieved using the perforated-patch clamp technique (Moser and Beutner, 2000) using Sylgard™-coated 1.5 mm borosilicate pipettes with typical resistances between 3.5 and 6 MΩ. The IHC pipette solution contained: 129 mM Cs-gluconate, 10 mM tetraethylammonium (TEA)-Cl, 10 mM 4-AP, 10 mM HEPES, 1 mM MgCl₂ (pH 7.2, osmolarity of ~290 mOsm), as well as 300 µg/ml amphotericin B added prior to the experiment. Once the series resistance of the IHC reached below 30 MΩ, whole-cell voltage-clamp recordings from a contacting bouton were established as described in previous studies (Glowatzki and Fuchs, 2002; Grant et al., 2010; Huang and Moser, 2018). Sylgard™-coated 1.0 mm borosilicate pipettes with typical resistances between 7 and 12 MΩ were used for the postsynaptic recordings. The bouton pipette solution contained: 137 mM KCl, 5 mM EGTA, 5 mM HEPES, 1 mM Na₂-GTP, 2.5 mM Na₂-ATP, 3.5 mM MgCl₂·6H₂O and 0.1 mM CaCl₂ (pH 7.2 and osmolarity of ~290 mOsm). The organ of Corti was continuously perfused with an extracellular solution containing 4.2 mM KCl, 95–100 mM NaCl, 25 mM NaHCO₃, 30 mM TEA-Cl, 1 mM Na-Pyruvate, 0.7 mM NH₂PO₄·H₂O, 1 mM CsCl, 1 mM MgCl₂·H₂O, 1.3 mM CaCl₂, and 11.1 mM D-glucose (pH 7.3, osmolarity of ~310 mOsm). 2.5 µM tetrodotoxin (Tocris or Santa Cruz) was added to block voltage-gated Na⁺ channels in the postsynaptic bouton.

Data were acquired using the Patchmaster software (HEKA electronics). The current signal was filtered at 5–10 kHz and sampled at 20–50 kHz. IHCs were voltage-clamped at a holding potential of –58 mV, corresponding to the presumed *in vivo* resting potential (Johnson, 2015). The bouton was held at a potential of –94 mV. All reported potentials were corrected for the liquid junction potential (19 mV for the IHC and 4 mV for the bouton), measured experimentally. Ca²⁺ current recordings were corrected for the linear leak current using a *P/n* protocol. We excluded IHCs and boutons with leak currents exceeding –60 pA and –100 pA at holding potential, respectively. The series resistance of the IHCs was typically below 30 MΩ. The apparent series resistance of the bouton was calculated from the capacitive transient in response to a 10-mV test pulse. The actual series resistance (*R_s*) was offline calculated as reported in (Huang and Moser, 2018). Recordings with bouton *R_s* > 80 MΩ were discarded.

The threshold for sEPSC detection was 4 times SD of the baseline. Spontaneous activity was calculated from time windows without stimulation with the IHC held at –58 mV; either from a 5–10 s recording or by averaging the number of events from the segments before and after a depolarizing pulse (Fig. 1B, Supplementary Table 1). To study the depletion and recovery of the pool of vesicles, we used a protocol adapted from the forward masking protocol performed during *in vivo* extracellular recordings of SGNs (Harris and Dallos, 1979; Jean et al., 2018). It consisted of two consecutive depolarizing pulses to the voltage that elicited the highest peak of Ca²⁺ current (–18 mV; Fig. 1C). The first pulse, called masker, lasted 100 ms and it was followed by a second pulse, called probe, which lasted 15 ms. The two pulses were separated by intervals without depolarization (interstimulus intervals, ISI) that lasted 4, 16, 64 and 256 ms. The waiting time between masker and masker was 20 s and each protocol was repeated between 3–20 times. To study the dynamic voltage range of synaptic transmission, we used a current-voltage (IV) protocol with 10 ms pulses of increasing voltage (from –70 mV/–60 mV to 70 mV in 5 mV steps). The interval between two stimuli was 1.5 s.

Data Analysis

Electrophysiological data was analyzed using the IgorPro 6 Software Package (Wavemetrics), GraphPad Prism 9 and Excel. Ca^{2+} charge (Q_{Ca}) and EPSC charge (Q_{EPSC}) were estimated by taking the integral of the current. Kinetics of sEPSCs, such as amplitude, 10-90% rise time, time constant of decay (τ_{decay}) and full-width half-maximum (FWHM), were calculated with Neuromatic (Rothman and Silver, 2018 [DOI](#)).

To obtain IV curves, we averaged the evoked Ca^{2+} -currents (I_{Ca}) during 5 to 10 ms after the start of each depolarization. Fractional activation of the Ca^{2+} channels was obtained from the normalized chord conductance, g ,

$$g = \frac{I}{(V - V_{\text{rev}})}$$

where V is the membrane potential and V_{rev} is the reversal potential determined by fitting a line function between the voltage of $I_{\text{Ca peak}} + 10$ mV and the maximal depolarization. The activation curve was approximated by a first-order Boltzmann equation:

$$g = \frac{g_{\text{max}}}{1 + \exp\left(\frac{V_{\text{half Ca}} - V}{S}\right)}$$

where g_{max} is the maximum chord conductance, $V_{\text{half Ca}}$ is the membrane potential at which the conductance is half activated, and S is the slope factor describing the voltage sensitivity of activation.

Release intensity curves were obtained by calculating Q_{EPSC} by the end of each depolarization step and fitted using a sigmoidal function:

$$Q = \frac{Q_{\text{max}}}{1 + \exp\left(\frac{V_{\text{h EPSC}} - V}{\text{rate}}\right)}$$

where Q_{max} is the maximal Q_{EPSC} (normalized to 1), $V_{\text{h EPSC}}$ corresponds to the voltage of half-maximal release (or $Q_{50, \text{EPSC}}$) and Q is the EPSC charge. The dynamic range was determined as the voltage range between 10% and 90% of the maximal vesicle release. For statistical analysis of dynamic range, we included only pairs for which both the Ca^{2+} fractional activation and the rate level curves were possible to fit.

The apparent Ca^{2+} dependence of neurotransmitter release was studied from the 10 ms step-depolarizations of the IV curves. The resulting Q_{EPSC} vs IHC Q_{Ca} plots from each individual pair were fitted with a power function:

$$Q_{\text{EPSC}} = a(Q_{\text{Ca}})^m$$

Where m corresponds to the Ca^{2+} cooperativity. Some pairs showed a clear saturation of release at high IHC Q_{Ca} . In these cases, the fit was restricted to the datapoints before the plateau, which was determined by visual inspection. For the pooled data, the power function was fitted to the normalized Q_{EPSC} vs normalized Q_{Ca} . For the pairs with saturation of release, Q_{Ca} was normalized to a point before the plateau.

For forward masking experiments, the postsynaptic response was averaged for all the repetitions for each paired recording (between 3 and 20, depending on the stability of the pair). Single active zone pool dynamics were determined by fitting an exponential plus line function to the individual EPSC charge plots,

$$y_0 + A1 \left(1 - \exp\left\{-\frac{(x-x_0)}{\tau}\right\} \right) + (x - x_0)\text{slope}$$

where $A1$ is the amplitude of the exponential component, τ is the time constant of the exponential component. RRP size (in vesicles) was estimated from dividing $A1$ by the charge of the average sEPSC for each pair. Sustained exocytosis rate (in vesicles per s) was calculated from the slope of the linear function divided the charge of the average sEPSC. Individual recovery kinetics were determined from the ratio of probe and masker responses at 10 ms of the depolarization, with the ratio between masker and masker being 1. The recovery traces were fitted with a single exponential function from 16 ms to 20 000 ms to determine the time constant of RRP recovery.

Data was prepared for presentation using Adobe Illustrator. Skewness analysis, PCA and other statistical analysis were performed using GraphPad Prism 9. Statistical significance was assessed with unpaired t-test or non-parametric Mann-Whitney U test depending on the normal distribution and equality of variances of the data (Saphiro-Wilk test and F test). Data is expressed as mean $\pm$ sem. The box plots show 25th, 50th and 75th percentiles with the individual data points overlaid.

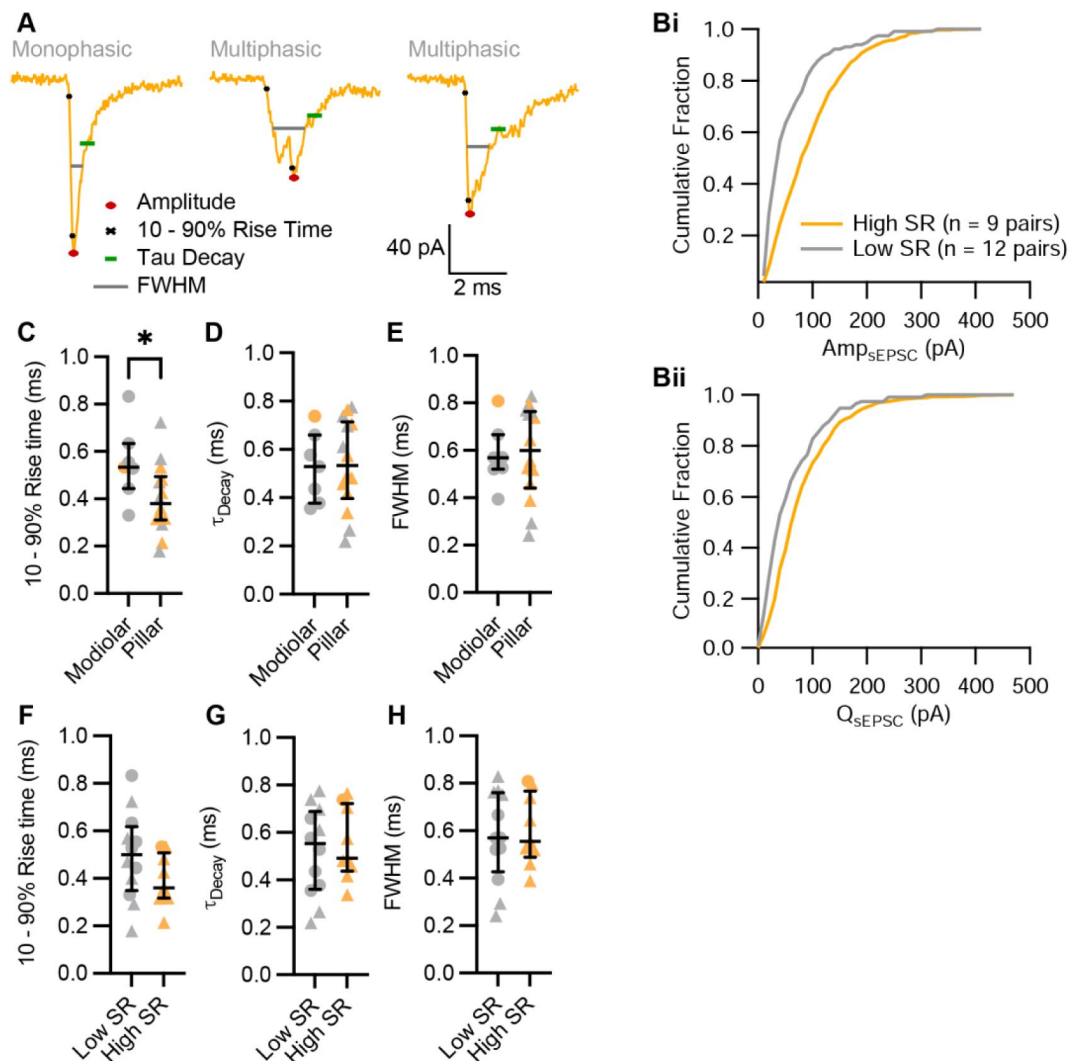
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Cellname	Side	SR	Total time for SR calculation	Cumulative time of 5-10 s recordings (s)	Cumulative time before/after a depolarizing pulse (s)
I140817_1	Modiolar	1.07	36		36
I140817_2	Modiolar	0.38	28.8		28.8
I140817_3	Modiolar	0.05	43.2		43.2
I300817_1	Modiolar	0.00	31.2		31.2
I240518_1	Pillar	0.22	91.2		91.2
I180618_1	Pillar	1.00	126.8		126.8
I031220_1	Pillar	0.00	10	10	
I041220_1	Pillar	0.00	10	10	
I280817_1	Modiolar	0.00	3.69		3.69
I200418_1	Pillar	3.92	3.06		3.06
I200618_1	Pillar	0.20	20.8		20.8
I260618_1	Pillar	2.05	20.8		20.8
I090718_2	Pillar	4.28	56.8		56.8
I170718_1	Pillar	0.38	23.2		23.2
I240718_4	Pillar	17.93	7.2		7.2
I250718_1	Pillar	3.18	38.4		38.4
I270718_1	Pillar	9.00	20		20
I090119_1	Modiolar	0.00	20	20	
I150618_1	Pillar	1.49	12		12
I290321_1	Modiolar	0.03	38	10	28
I300321_1	Modiolar	0.36	85.2	50	35.2
I120421_1	Pillar	0.21	35.6	10	25.6
I130421_1	Pillar	0.00	35.4	25	10.4
I301118_1	Modiolar	0.15	40	40	
I170119_1	Pillar	0.80	10	10	
I090419_1	Modiolar	0.00	10	10	
I120419_1	Modiolar	0.70	10	10	
I230920_2	Pillar	0.00	15	15	
I040221_1	Pillar	0.20	10	10	

Supplementary Table 1.

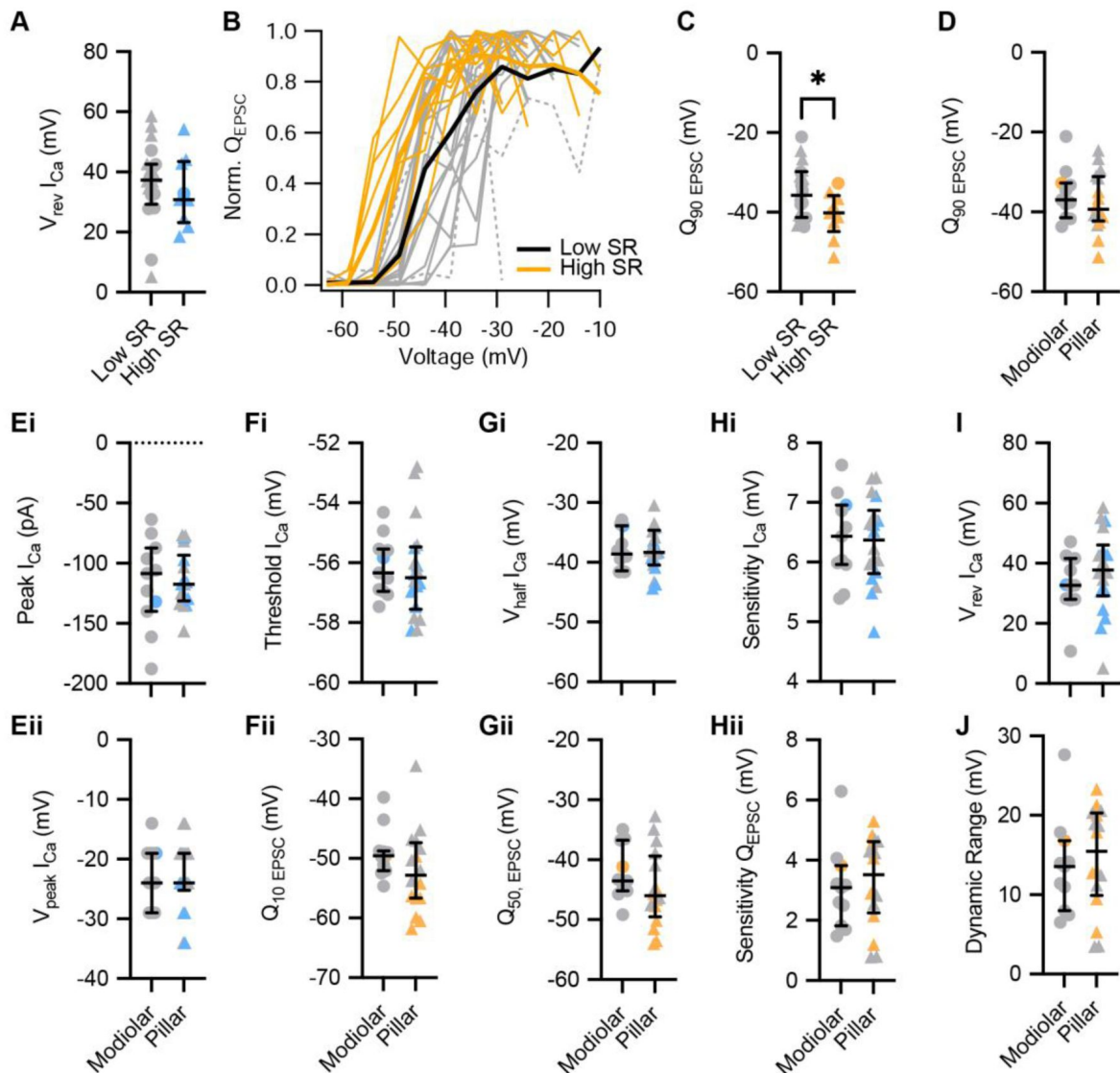
Spontaneous activity (SR) was calculated from time windows without stimulation with the IHC held at -58 mV (Total time for SR calculation). This total time was calculated from the cumulative recording time of either from 5 – 10 s recordings and/or from the segments before and after a depolarizing pulse.



Supplementary Figure 1.

The kinetics of sEPSCs are comparable between low and high SR synapses

A. Kinetics of sEPSCs, such as amplitude, 10-90% rise time, time constant of decay (τ_{decay}) and full-width half-maximum (FWHM), were calculated with Neuromatic (Rothman and Silver, 2018). **Bi-Bii.** Cumulative fraction of amplitude and charge of sEPSCs from low and high SR synapses. **C-E.** Pillar synapses had faster 10-90% rise times (C) than modiolar synapses ($p = 0.0426$, unpaired t-test), while their τ_{decay} (D) and FWHM (E) were comparable. **F-H.** Low and high SR synapses had similar 10-90% rise times (F), τ_{decay} (G) and FWHM (H). Scatter plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as $\triangle$ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.

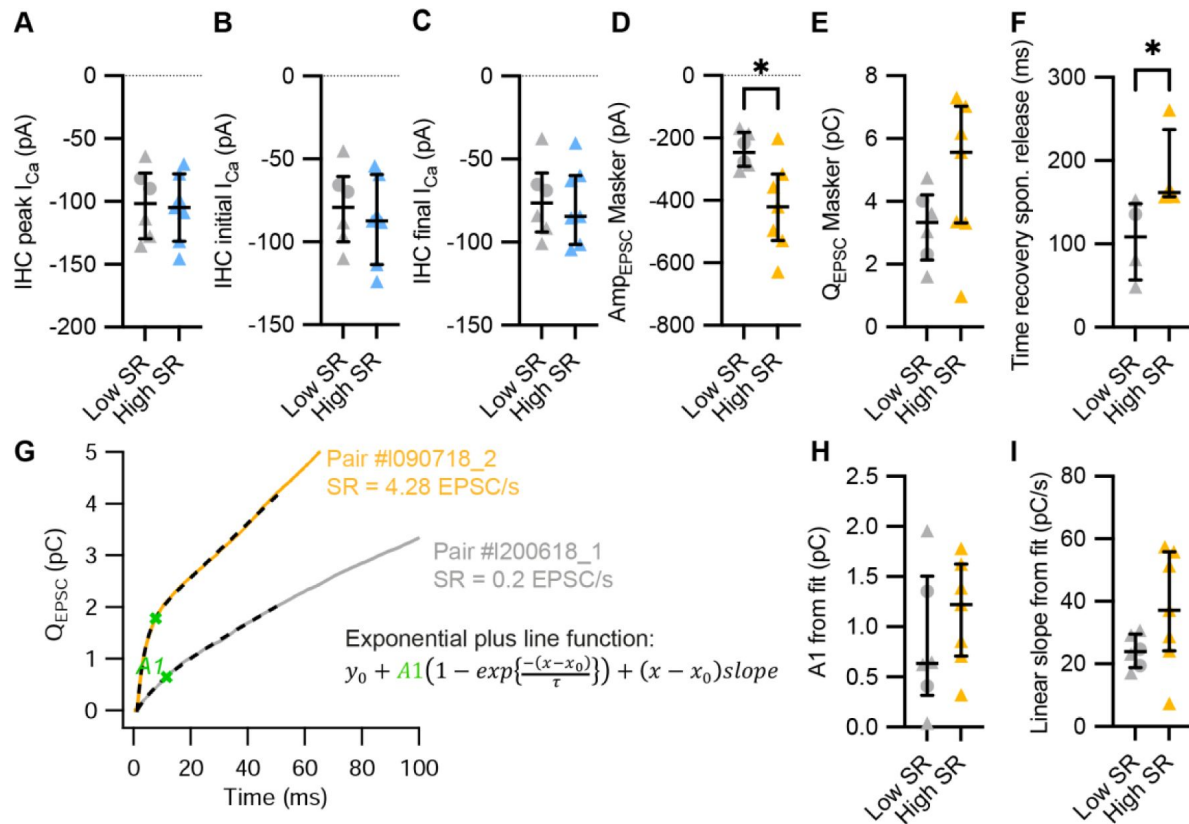


Supplementary Figure 2.

The voltage dependence of synaptic release does not differ between modiolar and pillar synapses

A. The reversal potential of Ca^{2+} influx was comparable between low and high SR synapses. **B.** Triggered single active zone eEPSCs (release intensity curves) of 29 pairs of low and high SR. Averages (thick lines) and individual curves (thin lines) are overlaid. The release-intensity curve of two pairs could not be fitted with the sigmoidal function (grey dotted lines). **C.** High SR synapses had more hyperpolarized voltages of 90% maximum release (Q_{90} EPSC, $p = 0.0396$, unpaired t test). **D.** Q_{90} EPSC did not differ between modiolar and pillar synapses. **Ei-Eii.** The peak of Ca^{2+} influx (Ei) and the voltage eliciting maximum Ca^{2+} current (Eii) were comparable between modiolar and pillar synapses. **Fi-Hi.** The threshold of Ca^{2+} influx (Ei), $V_{\text{half}} I_{\text{Ca}}$ (Fi) and voltage sensitivity of I_{Ca} (Gi) did not differ between modiolar and pillar synapses. **Fii-Hii.** Q_{10} EPSC (Fii), $V_{\text{half}} Q_{\text{EPSC}}$ (Gii) and voltage sensitivity of the release were comparable between modiolar and pillar synapses. **I.** The reversal potential of Ca^{2+} influx was comparable between modiolar and pillar synapses. **J.** Dynamic range of release was comparable between modiolar and pillar synapse.

Scatter plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as $\triangle$ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR ≥ 1 sEPSC/s.



Supplementary Figure 3.

Voltage dependence of IHC Ca^{2+} influx and synaptic release

A-C. Presynaptic peak (A), initial (B) and final (C) Ca^{2+} current (I_{Ca}) during the masker stimuli were comparable regardless the SR of postsynaptic bouton. **D-E.** High SR synapses had significantly larger amplitudes of evoked EPSCs in response to the masker stimulus (D; $p = 0.0147$, unpaired t-test), while the EPSC charge (Q_{EPSC}) did not differ (E). **F.** The spontaneous release after the probe offset recovered slower in high SR synapses. **G.** Exemplary average Q_{EPSC} response of a high SR (orange) and a low SR (gray) synapse to the masker stimuli. We fitted an exponential plus line function to the first 50 ms of the response (discontinuous lines) to study SV pool depletion dynamics. From these fits, we can retrieve information about RRP size (amplitude of the exponential component, A1), RRP depletion time constant (τ of the exponential component) and sustained release (linear slope of the line component). **H-I.** Amplitude (H) and linear slope (I) obtained from the fits of the exponential plus line function to the Q_{EPSC} from individual pairs (Fig. 4E). Scatter plots represent the 25th, 50th and 75th percentiles with the individual data points overlaid. Synapses were classified as $\triangle$ pillar or $\circ$ modiolar, and as Low SR < 1 sEPSC/s or High SR $\geq$ 1 sEPSC/s.

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Reviewer #1 (Public Review):

Summary:

Tobón and Moser reveal a remarkable amount of presynaptic diversity in the fundamental Ca dependent exocytosis of synaptic vesicles at the afferent fiber bouton synapse onto the pillar or mediolar sides of single inner hair cells of mice. These are landmark findings with profound implications for understanding acoustic signal encoding and presynaptic mechanisms of synaptic diversity at inner hair cell ribbon synapses. The paper will have an immediate and long-lasting impact in the field of auditory neuroscience.

Main findings: 1) Synaptic delays and jitter of masker responses are significantly shorter (synaptic delay: 1.19 ms) at high SR fibers (pillar) than at low SR fibers (mediolar; 2.57 ms). 2) Masked evoked EPSC are significantly larger in high SR than in low SR. 3) Quantal content and RRP size are 14 vesicles in both high and low SR fibers. 4) Depression is faster in high SR synapses suggesting they have a higher release probability and tighter Ca nanodomain coupling to docked vesicles. 5) Recovery of master-EPSCs from depletion is similar for high and low SR synapses, although there is a slightly faster rate for low SR synapses that have bigger synaptic ribbons, which is very interesting. 6) High SR synapses had larger and more compact (monophasic) sEPSCs, well suited to trigger rapidly and faithfully spikes. 7) High SR synapses exhibit lower voltage (~sound pressure in vivo) dependent thresholds of exocytosis.

Strengths:

Great care was taken to use physiological external pH buffers and physiological external Ca concentrations. Paired recordings were also performed at higher temperatures with IHCs at physiological resting membrane potentials and in more mature animals than previously done for paired recordings. This is extremely challenging because it becomes increasingly difficult to visualize bouton terminals when myelination becomes more prominent in the cochlear afferents. In addition, perforated patch recordings were used in the IHC to preserve its intracellular milieu intact and thus extend the viability of the IHCs. The experiments are tour-de-force and reveal several novel aspects of IHC ribbon synapses. The data set is rich and extensive. The analysis is detailed and compelling.

Weaknesses:

1. Materials and Methods: Please provide whole-cell R_s (series resistance) and C_m (membrane capacitance) average $\pm$ S.E.M. (or SD) values for IHC and afferent fiber bouton recordings. The C_m values for afferents have been estimated to be about 0.1 pF (Glowatzki and Fuchs, 2002) and it would be interesting to know if there are differences in these numbers for high and low SR afferents. Is it possible to estimate C_m from the capacitive transient time constant? Minimal electronic filtering would be required for that to work, so I realize the authors may not have this data and I also realize that the long cable of the afferents do not allow accurate C_m measurements, but some first order estimate would be very interesting to report, if possible.
2. Page 20, 26 and Figure 4: With regard to synaptic delays at auditory hair cell synapses: please see extensive studies done in Figure 11 of Chen and von Gersdorff (JNeurosci., 2019); this showed that synaptic delays are 1.26 ms in adult bullfrog auditory hair cells at 31°C, which is very similar to the High SR fibers (1.19 ms; Fig.4B and page 20). During ongoing depolarizations (e.g. during a sustained sine wave) the synaptic delay can be reduced to just 0.72 ms for probe EPSCs, which is a more usual number for mature fast synapses. This paper should, thus, be cited and briefly discussed in the Discussion. So a significant shortening of delay occurs for the probe response and this is also observed in young rat IHC synapses (see Goutman and Glowatzki, 2011).
3. Gaussian-like (and/or multi-peak) EPSC amplitude distributions were obtained in more mature rat IHCs by Grant et al. (see their Figure 4G; JNeurosci. 2010; postnatal day 19-21). The putative single quanta peak was at 50 pA and the main peak was at 375 pA. The large mean suggests a low CV (probably < 0.4). However, Fig. 2F shows a mean of about 100 pA and CV = 0.7 for spontaneous EPSCs. This major difference deserves some more discussion. I suppose that one possible explanation may be that the current paper holds the IHC membrane potential fixed at -58 mV, whereas Grant et al. (2010) did not control the IHC membrane potential and spontaneous fluctuations in the V_m may have depolarized the IHC, thus producing larger evoked EPSCs that are triggered by Ca channel openings. Some discussion that compares these differences and possible explanations would be quite useful for the readers.

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Reviewer #2 (Public Review):

Summary: The study by Jaime-Tobon & Moser is a truly major effort to bridge the gap between classical observations on how auditory neurons respond to sounds and the synaptic basis of these phenomena. The so-called spiral ganglion neurons (SGNs) are the primary auditory neurons connecting the brain with hair cells in the cochlea. They all respond to sounds increasing their firing rates, but also present multiple heterogeneities. For instance, some present a low threshold to sound intensity, whereas others have high threshold. This property inversely correlates with the spontaneous rate, i.e., the rate at which each neuron fires in the absence of any acoustic input. These characteristics, along with others, have been studied by many reports over the years. However, the mechanisms that allow the hair cells-SGN synapses to drive these behaviors are not fully understood.

Strengths:

The level of experimental complexity described in this manuscript is unparalleled, producing data that is hardly found elsewhere. The authors provide strong proof for heterogeneity in transmitter release thresholds at individual synapses and they do so in extremely complex experimental settings. In addition, the authors found other specific differences such as in synaptic latency and max EPSCs. A reasonable effort is put into bridging these observations with those extensively reported in *in vivo* SGNs recordings. Similarities are many and

differences are not particularly worrying as experimental conditions cannot be perfectly matched, despite the authors' efforts in minimizing them.

Weaknesses:

Some concern surges in relation to mismatches with previous reports of IHC-SGN synapses function. EPSCs at these synapses present a peculiar distribution of amplitudes, shapes, and rates. These characteristics are well-established and some do not seem to be paralleled in this study. Here, amplitude distributions are drastically shifted to smaller values, and rates of events are very low, all compared with previous evidence. The reasons for these discrepancies are unclear. The rate at which spontaneous EPSCs appear is an especially sensitive matter. A great part of the conclusions relies on the definition of which of the SGNs (or should say synapses) belong to the low end and which to the high end in the spectrum of spontaneous rates. The data presented by the authors seem a bit off and the criteria used to classify recordings are not well justified. The authors should clarify the origin of these differences since they do not seem to come from obvious reasons such as animal ages, recording techniques, mouse strain, or even species.

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Reviewer #3 (Public Review):

Summary:

"Bridging the gap between presynaptic hair cell function and neural sound encoding" by Jaime Tobon and Moser uses patch-clamp electrophysiology in cochlear preparations to probe the pre- and post-synaptic specializations that give rise to the diverse activity of spiral ganglion afferent neurons (SGN). The experiments are quite an achievement! They use paired recordings from pre-synaptic cochlear inner hair cells (IHC) that allow precise control of voltage and therefore calcium influx, with post-synaptic recordings from type I SGN boutons directly opposed to the IHC for both presynaptic control of membrane voltage and post-synaptic measurement of synaptic function with great temporal resolution.

Strengths

Any of these techniques by themselves are challenging, but the authors do them in pairs, at physiological temperatures, and in hearing animals, all of which combined make these experiments a real tour de force. The data is carefully analyzed and presented, and the results are convincing. In particular, the authors demonstrate that post-synaptic features that contribute to the spontaneous rate (SR) of predominantly monophasic post-synaptic currents (PSCs), shorter EPSC latency, and higher PSC rates are directly paired with pre-synaptic features such as a lower IHC voltage activation and tighter calcium channel coupling for release to give a higher probability of release and subsequent increase in synaptic depression. Importantly, IHCs paired with Low and High SR afferent fibers had the same total calcium currents, indicating that the same IHC can connect to both low and high SR fibers. These fibers also followed expected organizational patterns, with high SR fibers primarily contacting the pillar IHC face and low SR fibers primarily contacting the modiolar face. The authors also use in vivo-like stimulation paradigms to show different RRP and release dynamics that are similar to results from SGN in vivo recordings. Overall, this work systematically examines many features giving rise to specializations and diversity of SGN neurons.

Weaknesses / Comments / edits:

1. The careful analysis of calcium coupling and EPSC metrics is especially nice. Can the authors speculate as to why different synapses (likely in the same IHC) would have different calcium cooperativity?

2. On the bottom of page 6 it would be helpful to mention earlier how many pillar vs modiolar fibers were recorded from, otherwise the skewness of SRs (figure 2H could be thought to be due to predominantly recordings from modiolar fibers. As is, it reads a bit like a cliff-hanger.
3. The contrasts for some of the data could be used to point out that while significant differences occur between low and high SR fibers, some of these differences are no longer apparent when comparing modiolar vs pillar fibers (eg by contrasting Figure 2C and 2K). This can indicate that indeed there are differences between the fiber activity, but that the activity likely exists in a gradient across the hair cell faces. Pointing this out at the top of page 10 (end of the first paragraph) would be helpful, it would make the seemingly contradictory voltage-dependence data easier to understand on first read (voltage-dependence of release is significantly different between different SR fibers (figure 3) but is not significantly different between fibers on different HC faces (figure S3).
4. It should be acknowledged that although the use of post-hearing animals here (P14-23) ensures that SGN have begun to develop more mature activity patterns (Grant et al 2010), the features of the synapses and SGN activity may not be completely mature (Wu et al 2016 PMID: 27733610). Could this explain some of the 'challenges' (authors' section title) detailed on page 28, first full paragraph?
5. In the discussion on page 24, the authors compare their recorded SR of EPSCs to measure values in vivo which are higher. Could this indicate that in vivo, the resting membrane potential of IHCs is more depolarized than is currently used for in vitro cochlear experiments?
6. The results showing lower calcium cooperativity of high SR fibers are powerful, but do the authors have an explanation for why the calcium cooperativity of < 2 is different from that ($m = 3-4$) observed in other manuscripts?

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