Ca2+-permeable AMPARs mediate glutamatergic transmission and excitotoxic damage at the hair cell ribbon synapse

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Ca$^{2+}$-Permeable AMPARs Mediate Glutamatergic Transmission and Excitotoxic Damage at the Hair Cell Ribbon Synapse

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We report functional and structural evidence for GluA2-lacking Ca$^{2+}$-permeable AMPARs (CP-AMPARs) at the mature hair cell ribbon synapse. By using the methodological advantages of three species (of either sex), we demonstrate that CP-AMPARs are present at the hair cell synapse in an evolutionarily conserved manner. Via a combination of in vivo electrophysiological and Ca$^{2+}$ imaging approaches in the larval zebrafish, we show that hair cell stimulation leads to robust Ca$^{2+}$ influx into afferent terminals. Prolonged application of AMPA caused loss of afferent terminal responsiveness, whereas blocking CP-AMPARs protects terminals from excitotoxic swelling. Immunohistochemical analysis of AMPAR subunits in mature rat cochlea show regions within synapses lacking the GluA2 subunit. Paired recordings from adult bullfrog auditory synapses demonstrate that CP-AMPARs mediate a major component of glutamatergic transmission. Together, our results support the importance of CP-AMPARs in mediating transmission at the hair cell ribbon synapse. Further, excess Ca$^{2+}$ entry via CP-AMPARs may underlie afferent terminal damage following excitotoxic challenge, suggesting that limiting Ca$^{2+}$ levels in the afferent terminal may protect against cochlear synaptopathy associated with hearing loss.

Key words: cochlear synaptopathy; excitotoxicity; GCaMP; noise overexposure; zebrafish

Significance Statement

A single incidence of noise overexposure causes damage at the hair cell synapse that later leads to neurodegeneration and exacerbates age-related hearing loss. A first step toward understanding cochlear neurodegeneration is to identify the cause of initial excitotoxic damage to the postsynaptic neuron. Using a combination of immunohistochemical, electrophysiological, and Ca$^{2+}$ imaging approaches in evolutionarily divergent species, we demonstrate that Ca$^{2+}$-permeable AMPARs (CP-AMPARs) mediate glutamatergic transmission at the adult auditory hair cell synapse. Overexcitation of the terminal causes Ca$^{2+}$ accumulation and swelling that can be prevented by blocking CP-AMPARs. We demonstrate that CP-AMPARs mediate transmission at this first-order sensory synapse and that limiting Ca$^{2+}$ accumulation in the terminal may protect against hearing loss.

Introduction

Long-term studies of the mammalian cochlea have recently revealed that even a single incidence of noise overexposure can produce cochlear synaptopathy and neurodegeneration, exacerbating age-related hearing loss (Kujawa and Liberman, 2009; Lin et al., 2011). A major advance toward understanding this synaptopathy is to determine the cause of the initial excitotoxic insult.
This requires a more comprehensive understanding of the glutamate receptors that mediate synaptic function.

Much of our knowledge regarding the hair cell afferent synapse has been acquired from experiments performed in explanted endorgans of the prehearing animal (Glowatzki and Fuchs, 2002; Goutman and Glowatzki, 2007; Zhang-Hooks et al., 2016). It is widely assumed that glutamatergic transmission at this sensory synapse is mediated exclusively by Ca2+-impermeable AMPA receptors that include the Ca2+-limiting GluA2 subunit (Jonas and Burnashev, 1995). This assumption is supported by histological studies showing the expression of GluA2/3 and GluA4 at developing and mature cochlear hair cell synapses (Kuriyama et al., 1994; Matsubara et al., 1996; Fujikawa et al., 2014). Until now, there has been no evidence for postsynaptic regions lacking the GluA2 subunit and functional Ca2+-permeable AMPA receptors (CP-AMPARs) at the adult hair cell synapse.

To examine AMPARs that mediate synaptic transmission in maturing and adult synapses, we made use of the unique advantages of the following three model systems: zebrafish, rat, and bullfrog. The zebrafish skin is lined with clusters of hair cells that are both structurally and functionally similar to hair cells of the inner ear (Trapani and Nicolson, 2011; Maeda et al., 2014). Lateral line hair cells start mechanotransducing by 3 d postfertilization (Kindt et al., 2012), and by 5 dpf larvae are already using the lateral line to navigate their environment (Suli et al., 2012; Stewart et al., 2013). Given their external location, lateral line hair cells are optimal for in vivo Ca2+-imaging experiments and easily accessible for mechanical stimulation. Despite the methodological advantages and valuable insights gained from zebrafish (Söllner et al., 2004; Nicolson, 2005), novel findings in fish and other simple organisms are accompanied by questions regarding the relevance of new data to more complex organisms. To address this concern as well as to determine the applicability of lateral line findings to auditory organs, we performed work in both rat and frog auditory synapses. Functional studies, including Ca2+ imaging and electrophysiology, of the mature mammalian inner hair cell synapse are challenging given the limited access to hair cells and spiral ganglion neurons (SGNs; Chang et al., 2010; Wang et al., 2013). We thus made use of high-resolution immunohistochemical analysis of mature rat cochlear synapses to determine whether AMPAR subunit compositions that are consistent with CP-AMPARs can be observed. Last, we used the adult bullfrog preparation to determine whether CP-AMPARs mediate transmission in a bona fide auditory synapse. Each model organism provides a unique contribution to the work that addresses the caveats of the others to contribute receptors that mediate calcium accumulation in the auditory synapse.

In addition to examining synaptic transmission, we used zebrafish to examine the role of CP-AMPARs in excitotoxic damage of postsynaptic neurons. Calcium entry through ionotropic glutamate receptors (iGlurRs), primarily via NMDA receptors (NMDARs), underlies excitotoxicity throughout the nervous system (Lau and Tymianski, 2010). In some contexts, CP-AMPARs are the critical iGlurRs in this process (Liu and Zukin, 2007). In fact, the most promising evidence that CP-AMPARs may be expressed by SGNs comes from in vitro studies of AMPA-mediated excitotoxic damage in which AMPA exposure causes Ca2+ entry into afferent neurons (Eybalin et al., 2004).

In the present work, structural and functional studies of mature rat and frog auditory synapses were conducted to demonstrate that CP-AMPARs are present and mediate glutamatergic transmission in the auditory organ. Calcium-imaging studies in the larval zebrafish provided mechanistic information linking CP-AMPARs with excitotoxic damage and suggest a strategy for protecting against hearing loss by limiting Ca2+ accumulation in the afferent terminal.

**Materials and Methods**

**Zebrafish.** All zebrafish experiments were approved by the University of Washington Institutional Animal Care and Use Committee. Fish were reared in E3 embryo medium containing the following (in mM): 14.97 NaCl, 0.5 KCl, 0.042 NaHPO4, 0.15 KH2PO4, 1 CaCl2, 1 MgSO4, and 0.714 NaHCO3, pH 7.2, at ~28.5°C. Experiments were performed on zebrafish larvae 6–7 dpf, a stage before sex is determined in this species, in extracellular solution containing the following (in mM): 130 NaCl, 2 KCl, 1 MgCl2, 10 HEPES, and 2 CaCl2 (Trapani and Nicolson, 2011). Fish were paralysed via cardiac injection of α-bungarotoxin (125 µM; Trapani and Nicolson, 2010), which blocks skeletal muscle contraction while allowing the heart to continue beating. Briefly, fish were anesthetized by hypothermia, immobilized with a harp conventionally used for brain slices, and injected with α-bungarotoxin containing phenol red into the heart chamber. In a subset of imaging studies, fish were instead immobilized with 0.2% tricaine in E3. Extensive control experiments were conducted to determine whether tricaine affected waterjet responses, terminal morphology, and Ca2+ accumulation in the terminal. There were no differences between immobilization methods. When tricaine was used, it is indicated in the text.

**Transgenic zebrafish lines.** To assess hair cell and afferent terminal calcium responses to waterjet stimulation, Tg[myoD:GCaMP3]11001 (Estergberg et al., 2013) and Tg[elavl3:GCaMP5G]11001 (Ahrens et al., 2013) transgenic lines were used, respectively. To assess Ca2+ accumulation in afferent terminals, we used the Tg[elavl3:GCaMP5G] line. Given that elavl3 is a pan-neuronal promoter, we also generated zebrafish that transiently expressed GCaMP3 only in posterior lateral line ganglion (pLLG) afferent neurons driven by the SILLI enhancer [bsp70bGCaMP3-2.0ctnnap2a] (Pujol-Marti et al., 2012; see Fig. 9). To assess afferent terminal morphology, Tg[neuroD:eGFP]11001 (Obholzer et al., 2008) and Tg[islet1:GFP]11001 (Higashijima et al., 2000) transgenic lines were used, respectively.

**Zebrafish electrophysiology.** Paralyzed Tg[neuroD:eGFP] larvae were secured onto a Sylgard-coated coverslip with minimal application of WormGlu (GluStitch), and the fish and coverslip were transferred to a glass-bottom recording chamber coated with paraffin. Electrodes were filled with filtered extracellular solution. Loose-patch voltage-clamp recordings (series resistances, 20–80 MΩ) were obtained from individual neurons of the posterior lateral line ganglion visually identified using infrared differential interference contrast optics (Trapani and Nicolson, 2010). Data were acquired using pClamp10 software (Molecular Devices, 2005), and data were discarded if the resistance changed by >25%. Recordings were converted to text format then imported to MATLAB. Spike detection software in the Signal Processing Toolbox was used to quantify both spike frequency and interspike intervals.

**Waterjet stimulation.** Stereocilia of lateral line hair cells were displaced via application of a sinusoidal pressure wave (Trapani and Nicolson, 2011). A glass pipette (tip outer diameter, ~30 µm) was filled with an extracellular solution placed ~100 µm from the hair cell cluster termed a “neuromast,” and deflection of the kinocilia was used to verify pressure wave administration. Pressure output was driven by a pressure clamp (HSPC-1, ALA Scientific) that was commanded by pClamp10 software. To assess hair cell or afferent terminal responses to hair cell activation, a 20 or 10 s, respectively, rectified 1 Hz pressure wave was applied. GCaMP fluorescence values were captured every 1–2 s for 1 min using Slidebook with the onset of waterjet stimulation 20 s into capture.

**Pharmacology.** AMPA, cyclothiazide (CTZ), and α-bungarotoxin (Tocris Bioscience) were used at the indicated concentrations. To examine the receptors and pathways that mediate calcium accumulation in the terminals, we used antagonists for AMPARs (DNQX, Sigma Aldrich), Ca2+-permeable AMPARs [IEM1460 (herein referred to as IEM); Tocris Bioscience], and NMDARs (APV, Sigma Aldrich).
Zebrafish time-lapse imaging and analysis. Images were acquired using an inverted Marianas spinning disk system (Intelligent Imaging Innovations) equipped with an Evolve 10 MHz EMCCD camera (Photometrics) and a Zeiss C-Apochromat 63×/1.2 numerical aperture (NA) water-objective. Camera intensification was set to keep exposure times to <100 ms for GFP and ~100 ms for GCaMP, while keeping pixel intensity at <25% of saturation. Camera gain was set at 3 to minimize photobleaching. GFP and GCaMP fluorescence were acquired with a 488 nm laser and a 535/30 emission filter.

Given that all in vivo imaging studies were conducted on an inverted microscope, fish were oriented in two different ways for experiments in which we assessed terminal structure and Ca2+ accumulation versus waterjet responses. Fish were oriented with the dorsal side toward the microscope for all waterjet experiments. This enabled stimulation of the hair cell with the waterjet pipette and simultaneous imaging of the terminals. Terminals of P1–3 neuromast terminals were imaged from the side, and the D1 neuromast terminal was captured from the top down due to its more dorsal location. To assay terminal structure and quantify Ca2+ accumulation, fish were placed with their sides flush against the objective to visualize the organization of the terminal structure.

To examine morphological changes or Ca2+ levels in afferent terminals in response to AMPA application, Tg[neuroD:GFP] or Tg[elavl3:GCaMP5G] fish were immobilized with a harp. z-Sections were taken at intervals of 0.5–1 μm through the depth of the neuromast, including the afferent terminal, typically 20 μm. The z-sections were acquired before treatment, AMPA or DMSO was added directly to the imaging chamber, and images were acquired after 15 min of treatment. AMPA or DMSO was washed out of the chamber via a gravity perfusion system, and z-sections were again acquired 90 min postwash. Maximum projections in the untreated, AMPA, or DMSO condition, and if relevant, the wash condition were aligned using the Fiji StackReg plugin.

To monitor morphological changes in efferent terminals, the process was repeated with 6–7dpf Tg[Islet1:GFP] fish without a wash and recovery period. For analyses of morphological change, z-sections of Tg[neuroD:GFP]- or Tg[elavl3:GFP]-labeled processes were opened in the Fiji Simple Neurite Tracer (SNT). Neuritic processes of afferent and efferent terminals were traced for each of the treatment conditions (afferent: control, AMPA, and wash; efferent: control and wash). Using the maximum projection image for each condition, processes were traced starting from the first branch point along the axon to the most distal point that likely represented a postsynaptic terminal. Each neurite was traced from a branch point to the distal most pixel identified by SNT. SNT only traced neurites represented by a continuous segment of bright pixels. Terminal swelling often caused breaks in this continuous segment or shortening of the neuritic branch. All values were converted to microns using the appropriate calibration for the microscope settings. Mean branch length and number of branches were computed for each terminal in the treatment conditions.

To assess hair cell and afferent terminal calcium responses to waterjet stimulation, hair cells or terminals were identified that consistently responded to at least two stimulation trials. AMPA was applied to the imaging chamber for at least 15 min, and two stimulation trials were repeated. For all hair cell-imaging experiments, a single hair cell was analyzed for each Tg[myo6b:GCaMP3] fish (n = 7 fish). The region of interest was drawn around the entire hair cell that showed a response. For all terminal imaging, a single terminal was imaged in each Tg[elavl3:GCaMP5G] fish, and the entire region of the terminal shown in the focal plane was designated as the region of interest. To assay the recovery of afferent terminal responses, AMPA was washed out of the imaging chamber for at least 60 min, and the stimulation trials were repeated.

For analyses of waterjet hair cell and terminal responses, fluorescence intensity was quantified using SlideBook software (Intelligent Imaging Innovations) and GCaMP fluorescence measurements were exported to MATLAB. As described previously (Esterberg et al., 2013), fluorescence data are normalized by dividing the fluorescence intensity values for the entire acquisition period by the baseline value before waterjet stimulation. Using baseline-adjusted traces, the peak of each calcium response was computed. To compute the half-width of each trace as a measure of response duration, traces were first normalized to peak. Each data point in the control, AMPA-treated, and recovery conditions represents the mean of two GCaMP responses to a pressure wave.

Zebrafish whole-mount immunohistochemistry and fluorescence imaging. All zebrafish immunohistochemistry experiments were approved by the Massachusetts Eye and Ear Institution Animal Care and Use Committee (IACUC). The 5-d-old larvae were exposed to 100 μM (S)-AMPA (Abcam) diluted in E3 embryo media with 0.1% DMSO for 15 min at room temperature (20–22°C). E3 with 0.1% DMSO were used as controls. Following drug exposure, larvae either were immediately fixed for immunohistochemistry or were allowed to recover for 1 h. Larvae were quickly sedated on ice then transferred to fixative (4% paraformaldehyde/4% sucrose/0.15 mM CaCl2 in phosphate buffer) for 1 h. Following fixation, larvae were permeabilized in 1% Tween in PBS for 5 h at 4°C then blocked in PBS buffer containing 1% bovine serum albumin, 0.5% Fish Skin Gelatin, 2% goat serum, and 0.1% DMSO for >3 h. The following antibodies were used: MAGUK (K28/86; 1:500); GluA4 (1:400; Millipore Bioscience Research Reagents); and an affinity-purified antibody generated against zebrafish Ribeye b (mouse IgG2a; 1:2000; Sheets et al., 2011). Primary antibodies were diluted in blocking buffer and applied overnight at 4°C followed by diluted secondary antibodies coupled to Alexa Fluor 488 and Alexa Fluor 647 (Invitrogen), or DyLight 549 (Jackson ImmunoResearch), and labeled with DAPI (Invitrogen). z-Stack images of neuromasts (spaced by 0.3 μm over 10–12 μm) were acquired with a Leica SP8 confocal microscope using a 63×/1.3 NA glycerol immersion objective lens. For each experiment, the microscope parameters were adjusted using the brightest control specimen. Digital images were processed using ImageJ software and Amira 3D software (Interactive3D).

Quantitative image analysis was performed on raw images using Amira 3D Analysis software as previously described (Lv et al., 2016). To quantitatively measure immunolabel intensity, a user-defined inclusive threshold was applied to isolate pixels occupied by GluA4 or postsynaptic density (PSD)-immunolabeled patches. The inclusive threshold values for each label were determined using 3D isosurface renderings, with the minimum threshold value defined as the value above which the user could resolve two closely adjacent patches. The material statistics function was then used to measure the cumulative intensity of fluorescent pixels (sum of the grayscale values) within each individual sphere or patch as well as to define the x-, y-, and z-coordinates of the center of each puncta within the stack. Postsynaptic Gria4-immunolabeled clusters were identified as Gria4 labeling that generally overlapped with the MAGUK immunolabel.

Rat Gill5 staining. All rat experiments were approved by the IACUC at Washington University in St. Louis. Immunohistochemistry on mature Wistar rats of either sex was performed as previously described (Jing et al., 2013; Kim and Rutherford, 2016). Cochlear whole-mount samples were washed in the PDS overnight and imaged in the organ of Corti on a Zeiss LSM 700 microscope with a 63× 1.4 NA oil-immersion objective. Pixels were 50 × 50 nm, and the z-step was 333 nm.

Three AMPA receptor subunits were labeled simultaneously with primary antibodies against GluA2 (mouse IgG2a, catalog #MAB397, Millipore; Vis-savajihala et al., 1996), GluA3 (goat, sc-7612, Santa Cruz Biotechnology), or GluA4 (rabbit, catalog #AB1508, Millipore; Wenthold et al., 1992). Specificity of the GluA2 and GluA4 antibodies (Medvedev et al., 2008; Sagata et al., 2010) were verified in GluA2 and GluA4 knock-out mice (Jia et al., 1996; Sagata et al., 2010), respectively. We find no reference for use of the GluA3 antibody on the GluA3 knock-out mouse; however, sc-7612 has been used previously with the GluA4 antibody AB1508 for immunohistochemistry on retinal ribbon synapses with similar results (Puller and Haverkamp, 2011; Puthussery et al., 2014). Secondary antibodies were Alexa Fluor 488 anti-goat, Alexa Fluor 555 anti-rabbit, and Alexa Fluor 647 anti-mouse. Imaris software (Bitplane) was used for 3D analysis of synaptic punctum intensity and volume. We note that differences in the levels of subunit immunofluorescence potentially result from differences in the binding efficiency of the antibodies, and thus acquiring three channels with the same microscope settings would not be expected to produce fluorescence ratios that indicate absolute measurements of subunit relative abundance at a given synapse. Therefore, to maximize the dynamic range of level detection during acquisition, the microscope settings on each of the three channels in an image.
volume were adjusted to have level saturation at the brightest pixel of the brightest synapse. This normalization resulted in synapses with similar overall intensities on each channel. IGOR software (WaveMetrics) was used for line plots on the 2D maximum-intensity projections of postsynaptic densities oriented en face. Images in Figure 11 were deconvolved, to reduce blur, with an optical model of the point spread function in ImageJ software as previously described (Jing et al., 2013; Ohn et al., 2016).

To control for artifacts due to bleaching and attenuation in our experimental system, we coated coverslips with fluorescent beads and mounted the tissue between two coverslips so that the sample and the beads on both sides can be imaged and measured from either side. Attenuation makes synapses closer to the coverslip appear brighter. Assuming a linear trend, we estimate the effects of attenuation in the whole mount to be <5% for synapses separated by <20 μm along the z-axis. Local deviations in volume and intensity between synapses (i.e., between synapses centered within 333 nm from each other) far exceed the size of the linear trend across the ~20 μm range of the synaptic layer, suggesting that true morphological heterogeneity dominates these differences in luminance values. Fluorophores outside the focal plane can be bleached, leading to artifacts that depend upon the order of acquisition when scanning through Z. To control for bleaching, we scanned the volume twice and calculated the rate of bleaching for each channel by comparing the synaptic intensities from scan 1 with the synaptic intensities from scan 2. For GluA2 per synapse, the Corrected Intensity/0.991^section#, where section# was calculated from the data for each subunit for each whole mount to be –6 fish, 1 terminal/fish). Paired t test.

Figure 1. Waterjet-mediated hair cell stimulation causes afferent terminal Ca2+ entry. Tg(elavl3:GCaMP5G) transgenic line was used to measure responsiveness of afferent terminal to waterjet-mediated displacement of hair cell stereocilia. A, B, Frames from a time-lapse video (1 s capture interval) before, during, and after waterjet stimulation are shown in the control condition (A) and following DMSO application (B). Dashed line marks the location of the terminal and thereby the region of interest used for analysis. Scale bar, 10 μm. C, Representative traces showing changes in the fluorescence intensity normalized to baseline. Horizontal line starting at time = 0 marks the duration of the waterjet stimulus. D, E, Summary data of mean peak and integrated area of the response in the control and DMSO condition (n = 6 fish, 1 terminal/fish). Paired t test.

Figure 2. Perfusion of CTZ enhances terminal responsiveness to hair cell activation. Tg(elavl3:GCaMP5G) transgenic line was used to measure the responsiveness of the afferent terminal to waterjet-mediated displacement of hair cell stereocilia. A, B, Maximum projections of an afferent terminal in the control condition (A) and with CTZ (B). C, D, Frames from a time-lapse video (1 s capture interval) before, during, and after waterjet stimulation are shown in the untreated condition (C) and following CTZ application (D). The dashed line marks the location of the terminal and thereby the region of interest used for analysis. Scale bar, 10 μm. E, Representative traces showing changes in the fluorescence intensity normalized to baseline. F, G, CTZ increases the peak and integrated area of the waterjet response. Paired t test.

Electrophysiology from adult bullfrog auditory hair cells. All frog experiments were approved by the IACUC at Oregon Health & Science University. Amphibian papillae of adult female or male bullfrogs (Rana catesbeiana) were carefully dissected and, as previously described (Keen and Hudspeth, 2006; Li et al., 2009), semi-intact preparations of hair cells and their corresponding afferent fibers were obtained. During the re-
cordings, the preparation was perfused (2–3 ml/min) with oxygenated artificial perilymph containing the following (in mM): 95 NaCl, 2 KCl, 2 CaCl₂, 1 MgCl₂, 25 NaHCO₃, 3 glucose, 1 creatine, and 1 Na-pyruvate, pH 7.30, 230 mOsm. For paired recordings, patch pipettes of borosilicate glass were pulled to resistances of 5–6 MΩ for hair cells and 7–10 MΩ for afferent fibers. Pipettes were filled with the internal solution containing the following (in mM): 77 Cs-gluconate, 20 CsCl, 1 MgCl₂, 10 TEA-Cl, 10 HEPES, 2 EGTA, 3 Mg-ATP, 1 Na-GTP, and 5 Na₂-phosphocreatine, pH 7.30, 230 mOsm. Whole-cell recordings were performed at room temperature using an EPC-10/2 amplifier (HEKA) and Patchmaster software (HEKA). Both hair cells and afferent fibers were voltage clamped with a resting membrane potential of ~90 mV after liquid junction potential correction. The current signal was low-pass filtered at 5 kHz and sampled at 10 μs intervals. The averaged uncompensated series resistances in whole-cell recordings were 12.6 ± 1.0 MΩ for hair cells (n = 16) and 23.8 ± 2.8 MΩ for afferent fibers (n = 16). The measurements of the whole-cell membrane capacitance (C_m) from hair cells were performed under voltage clamp with the “Sine + DC” method (Lindau and Neher, 1988; Gillis, 2000) using an EPC-10/2 patch-clamp amplifier (HEKA) and Pulse software (HEKA). Sine waves (50 mV peak-to-peak, 1 or 2 kHz) were superposed on the holding potential, and the resulting current response was used to calculate C_m via a Pulse software emulator of a lock-in amplifier (Gillis, 2000). Data analysis was performed with Igor Pro software (WaveMetrics). Data are expressed as the mean ± SEM.

Statistics. GraphPad Prism 5.0 Software was used for all statistical analyses. ANOVAs used the Tukey post hoc test.

Figure 3. AMPA perfusion diminishes afferent neuron firing in a dose-dependent manner. A, Schematic showing zebrafish head and the location of the pLLG just caudal to the ear. Loose-patch voltage-clamp recordings were obtained from cell bodies in the pLLG (green), which receive glutamatergic inputs from clusters of hair cells (red). B, Representative traces of pLLG neuron firing in ambient conditions (control) and following bath application of varying doses of AMPA. AMPA (100 and 300 μM) caused a statistically significant transient increase in firing frequency in many cells during the first 1–2 min of application (*p < 0.05) followed by a dramatic reduction in firing (#p < 0.001). C, Summary data show that the mean firing frequency decreases dramatically following 100 μM (n = 9 cells) and 300 μM (n = 6 cells) AMPA application but not 30 μM AMPA (n = 8 cells). Two-way ANOVA.

Figure 4. Perfusion of 100 μM AMPA diminishes terminal responsiveness to hair cell activation. The Tg(eln3:GCaMP5G) transgenic line was used to measure the responsiveness of afferent terminal to waterjet-mediated displacement of hair cell stereocilia. A–C, Frames from a time-lapse video before, during, and after waterjet stimulation are shown in the control condition (A), following 100 μM AMPA application (B), and 2 h after wash (C). The dashed line marks the location of the terminal and thereby the region of interest used for analysis. Scale bar, 10 μm.

D, Response to waterjet is diminished following AMPA application and wash. E, F, Summary data of the mean peak and integrated area of the response in the control, AMPA, and wash conditions. One-way ANOVA, *p < 0.05.
Results

In vivo hair cell stimulation causes Ca\(^{2+}\) influx into afferent terminals

To examine the contribution of AMPARs to glutamatergic transmission at the hair cell synapse, we used in vivo time-lapse Ca\(^{2+}\) imaging in the larval zebrafish lateral line. We visualized afferent terminal responses to hair cell stereocilia deflection using the Tg[elavl3:GCaMP5G] line in which the Ca\(^{2+}\) indicator GCaMP5 is expressed pan-neuronally, including in afferent neurons. Stereocilia were displaced for 20 s using a 1 Hz rectified pressure wave administered via a waterjet pipette placed ~100 \(\mu\)m away from the neuromast. Figure 1 shows frames from time-lapse videos with single planes of afferent terminals before, during, and after waterjet stimulation. Under control conditions, GCaMP5 fluorescence intensity increases, indicating that hair cell activation causes an influx of Ca\(^{2+}\) into the terminal (Fig. 1A, F). The response to waterjet stimulation was quantified by measuring the mean peak fluorescence and duration of the response across two waterjet stimulation trials.

AMPARs desensitize at a faster time course relative to NMDA and kainate receptors (Geiger et al., 1995), a process that can be studied using the drug CTZ, which blocks AMPAR desensitization (Trussell et al., 1993). Bath application of 200 \(\mu\)M CTZ had no detectable effect on GCaMP fluorescence under resting conditions, as shown in maximum projections of a \(z\)-stack of a terminal before and after CTZ application (Fig. 2A, B). In contrast, CTZ (200 \(\mu\)M) enhanced the terminal waterjet responses as shown in the time-lapse video frames (Fig. 2C, D) and the plot of fluorescence intensity over time (Fig. 2E). CTZ increases the mean peak (control, 1.12 ± 0.02; CTZ, 1.18 ± 0.02; \(n = 7\), \(p < 0.05\), mean ± SEM, paired \(t\) test; Fig. 2F) and integrated area (con-
trol, 2.6 ± 0.3; CTZ, 3.4 ± 0.5; n = 7, p < 0.05, mean ± SEM; Fig. 2G) of the waterjet responses.

We attempted to determine whether the terminal Ca\(^{2+}\) responses are mediated by CP-AMPARs using an antagonist that specifically blocks these receptors, IEM (Kim and von Gersdorff, 2016). We imaged waterjet responses before and after bath application of 100 μM IEM or DMSO carrier. DMSO does not substantially change the mean integrated area (control, 2.6 ± 1.0; DMSO, 3.1 ± 1.3; n = 6, p = 0.15, mean ± SEM) of the terminal response. In contrast, IEM significantly diminished terminal waterjet responses as shown in the time-lapse video frames and the plot of fluorescence intensity over time. IEM decreased the mean integrated area (control, 3.5 ± 0.6; IEM, 1.1 ± 0.09; n = 5, p < 0.05, mean ± SEM) of the waterjet responses. However, imaging of hair cell waterjet responses using Tg[myo6b:GCaMP3]\(^{w78}\) revealed that IEM (50 and 100 μM) also blocks hair cell responses (control, 4.5 ± 0.8; IEM, 1.3 ± 0.1; n = 6, p < 0.05, mean ± SEM). Fortunately, the bullfrog model system was better suited to examine the contribution of CP-AMPARs to glutamatergic transmission at the hair cell synapse (see Fig. 12). However, we continued using the zebrafish model system to examine the contribution of CP-AMPARs to excitotoxic damage.

Afferent neurons lose responsiveness after exposure to exogenous AMPA

We conducted a series of electrophysiological and Ca\(^{2+}\) imaging experiments to determine whether CP-AMPARs mediate excitotoxic damage. First, we examined the effect of AMPA application on the overall activity of posterior lateral line ganglion neurons by recording presynaptically driven spontaneous spike firing. The range of concentrations was chosen based on dose-response studies in mammalian cochlea (Puel et al., 1991, 1994). In Figure 3A is a schematic of a larval zebrafish head showing the location of the recording electrode used to obtain loose-patch voltage-clamp recordings from individual pLLG neurons. Figure 3B shows representative action current recordings before and after the perfusion of varying concentrations of AMPA. Application of 100 or 300 μM AMPA caused a brief bout of high-frequency spiking before firing diminished (Fig. 3B, middle panels). We observed the initial increase in spiking in six of nine neurons and the decrease in all neurons analyzed. The AMPA concentration of 100 μM was the lowest dose tested that reduced mean firing frequency (Fig. 3C).

We next examined how the response of afferent terminals to waterjet stimulus is altered by AMPA treatment. We imaged waterjet responses in a terminal in the untreated condition, following bath application of 100 μM AMPA and 2 h after wash (Fig. 4A–C). We noted that AMPA exposure increased the baseline fluorescence in terminals, examined further below. However, after AMPA exposure there was little or no increase in terminal fluorescence in response to waterjet. AMPA decreased the mean peak (Fig. 4E; control, 1.2 ± 0.5; AMPA, 1.1 ± 0.5; wash, 1.0 ± 0.5; n = 6, mean ± SEM, one-way ANOVA; control vs AMPA, p < 0.05) and integrated area (Fig. 4F; control, 2.2 ± 1.0; AMPA, 1.0 ± 0.5; wash, 1.0 ± 0.5; n = 6, mean ± SEM, one-way ANOVA; control vs AMPA, p < 0.05; control vs wash, p < 0.05) of the terminal response, and these values did not recover after wash.

While these results are consistent with AMPA overexposure resulting in postsynaptic dysfunction, it is possible that AMPA disrupted neuronal firing indirectly by acting on the presynaptic hair cell. To address whether AMPA exposure altered the hair cell response to mechanical stimuli, we conducted in vivo Ca\(^{2+}\)-imaging studies using the Tg[myo6b:GCaMP3]\(^{w78}\) fish, which express GCaMP3 in the hair cell cytoplasm during and after waterjet exposure (Fig. 5). Under control conditions, increases in fluorescence intensity temporally correlate with the duration of waterjet stimulation. AMPA did not substantially change either the mean peak (control, 1.6 ± 0.2; AMPA, 1.5 ± 0.1; n = 7, p = 0.9, mean ± SEM) or integrated area (control, 8.8 ± 2.6; AMPA, 8.9 ± 2.0; n = 7, p = 0.97, mean ± SEM) of the hair cell Ca\(^{2+}\) response to mechanical displacement, suggesting that presynaptic Ca\(^{2+}\) signals were maintained during AMPA exposure and that loss of responsiveness was mediated postsynaptically.

Exogenous AMPA exposure results in excitotoxic damage that involves CP-AMPARs

Our observation that AMPA exposure caused a loss of terminal responsiveness suggested that the terminal sustained excitotoxic damage. One characteristic of such damage in the mammalian cochlea is a swelling of afferent SGN terminals (Puel et al., 1991, 1994, 1998). We conducted in vivo time-lapse imaging of the Tg[neuroDx::GFP] transgenic line, which labels afferent terminals before, during application of 100 μM AMPA, and 90 min after washout (Fig. 6). During AMPA application, terminals exhibited swelling consistent with mammalian observations. In the panel of images shown in Figure 6A, these swellings are marked by the arrowheads in the AMPA treatment condition (Fig. 6A, middle). The number of neuritic branches in each terminal decreased during AMPA treatment (control, 9.9 ± 0.6; AMPA, 6.1 ± 0.6; wash, 6.0 ± 0.5; n = 21, mean ± SD; ANOVA, p < 0.0001) while the mean neuritic branch length remained the same (control, 13.8 ± 0.5 μm; AMPA, 15.5 ± 0.9 μm; wash, 15.9 ± 1.1 μm; n = 21, mean ± SEM). Following 90 min of washing, neither measure of terminal morphology recovered. Figure 6A–F represent data acquired in larvae immobilized with α-bungarotoxin versus tricaine. In the presence of tricaine (Fig. 6D–F), the number of branches decreased during AMPA treatment (control, 6.1 ± 0.5; AMPA, 4.5 ± 0.4; n = 16, mean ± SD; paired t test, p < 0.001) while the mean branch length remained the same (control, 15.7 ± 0.6 μm; AMPA, 17 ± 1.2 μm; n = 16, mean ± SEM). The immobilization method did not affect AMPA-mediated changes in terminal morphology. Blocking CP-AMPARs via bath application of IEM before 100 μM AMPA treatment prevented changes to the number...
of branches (control, 5.4 ± 0.4 μm; IEM, 6.1 ± 0.5 μm; IEM plus AMPA, 6.1 ± 0.4 μm; n = 18, mean ± SEM). IEM treatment alone slightly reduced the mean branch length relative to control (control, 15 ± 1 μm; IEM, 14 ± 1 μm; IEM plus AMPA, 14 ± 1 μm; n = 18, mean ± SEM; ANOVA, p < 0.05). Representative images (Fig. 6G) and summary data (see Fig. 9H, I) for the control, IEM, and IEM plus AMPA conditions are shown.

We also examined whether AMPA causes any gross morphological changes to the presynaptic efferent terminals that are thought to play an inhibitory modulatory role at this synapse. We monitored the morphology of efferent terminals using time-lapse imaging experiments of Tg[Islet1:GFP]. Maximum projections of an efferent terminal are shown in Figure 7A before and during 100 μM AMPA exposure. There were no changes in the number of branches (control, 8 ± 3; AMPA, 8 ± 3; n = 9, mean ± SD) or mean branch length (control, 10 ± 0.6 μm; AMPA, 10 ± 1 μm; n = 9, mean ± SEM) in the presence of AMPA (Fig. 7B, C), demonstrating that efferent synapses were relatively insensitive to AMPA exposure.

To determine whether the distribution of synapses was altered after AMPA over-exposure, we conducted triple immunolabeling of the presynaptic protein Ribeye, postsynaptic density MAGUK, and the GluA4 subunit (Fig. 8). Although gria2a and gria2b, encoding for GluA2 subunits, are expressed in the zebrafish lateral line ganglia (Hoppmann et al., 2008), commercial GluA2 antibodies do not cross-react with zebrafish. AMPA treatment had little effect on the expression of GluA4 nor did it change the distribution of Ribeye- and GluR-expressing puncta, suggesting that this AMPA exposure silenced synapses without their physical disassembly.

As noted earlier, we observed that terminals treated with AMPA exhibit an overall baseline increase in GCaMP fluorescence intensity while at the same time blocking response to water jet stimuli. We quantified the percentage change in GCaMP fluorescence intensity following bath application of increasing concentrations of AMPA, compared with untreated terminals or those treated with DMSO carrier alone. While the application of DMSO alone or with 10 or 30 μM AMPA had no effect on GCaMP fluorescence of terminals (Fig. 9A, B, I), treatment with 100 and 300 μM AMPA (Fig. 9C, D, I) resulted in robust increases in fluorescence intensity relative to the untreated condition.

To determine which glutamatergic receptors mediate Ca\(^{2+}\) accumulation, we quantified the GCaMP fluorescence levels after a 15 min application of 100 μM AMPA in the presence of glutamate receptor blockers. Pretreatment with 40 μM DNQX, an antagonist for the AMPA and kainate receptors, blocked the increase in terminal GCaMP fluorescence levels that are observed following AMPA exposure (Fig. 9E, I). In contrast, NMDAR blockade using 50 μM APV was not able to block AMPA-mediated accumulation of terminal Ca\(^{2+}\). We next examined whether blocking only CP-AMPARs can protect the terminal from Ca\(^{2+}\) accumulation and putative excitotoxic damage. At all doses tested, the CP-AMPAR blocker (IEM) was sufficient to either reduce or block AMPA-mediated increases in GCaMP fluorescence (Fig. 9H, K). To further demonstrate that AMPA causes Ca\(^{2+}\) accumulation in pLLG afferent neurons that is prevented by blocking CP-AMPARs, we generated transgenic zebrafish that transiently express GCaMP3 in pLLG neurons. AMPA (100 μM) caused a significant increase in fluorescence intensity that was prevented by blocking CP-AMPARs with IEM (100 μM) pretreatment (Fig. 9L–O).

We tested whether ion channels contributed to Ca\(^{2+}\) accumulation in afferent terminals. To test whether voltage-
gated Na\(^+\) and K\(^+\) channels are a requirement for Ca\(^{2+}\) accumulation, we anesthetized fish via bath application of the anesthetic tricaine. We found that Ca\(^{2+}\) accumulation was unchanged by tricaine. Moreover, AMPA had the same effects on GCaMP fluorescence regardless of whether fish were paralyzed with α-bungarotoxin or anesthetized with tricaine (Fig. 9); two-way ANOVA. Pretreatment with nifedipine, an L-type voltage-gated Ca\(^{2+}\) channel blocker, did not block Ca\(^{2+}\) accumulation; rather, it somewhat enhanced AMPA-mediated Ca\(^{2+}\) entry. Moreover, AMPA concentration relative to control and following application of one of the receptor antagonists showed that AMPA has a dose-dependent effect on the accumulation of terminal Ca\(^{2+}\). For all data points, the fluorescence intensity of a single terminal was measured in the untreated control condition and following drug application. The 100 and 300 μM concentrations of AMPA significantly increase the fluorescence intensity of terminals. This increase is blocked by DNQX and IEM, but not by the NMDAR blocker APV. I, AMPA has a dose-dependent effect on the accumulation of terminal Ca\(^{2+}\) as shown in the plots above the data points. When we applied 100 μM AMPA, the GCaMP fluorescence over AMPA alone (percentage change in terminal fluorescence intensity relative to the untreated control condition). J, Summary data showing that AMPA has a dose-dependent effect on the accumulation of terminal Ca\(^{2+}\) at all doses tested. L, Maximum projections of afferent terminals in the untreated condition (control) and following application of DMSO, AMPA alone or application of a IEM followed by 100 μM AMPA. O, Summary data showing that AMPA causes significant increases in fluorescence intensity that are blocked by IEM. Scale bars, 10 μm. One-way ANOVA: \(p < 0.05\) \(*\) \(p < 0.01\); \(**p < 0.001\); \(***p < 0.0001\). All values are indicated in all plots above the data points.

GluA2-lacking AMPARs reside at peripheral auditory synapses

To determine whether CP-AMPARs are found within the mature mammalian cochlea, we undertook quantitative analysis of the relative abundance of AMPAR subunits. Each mature mammalian auditory nerve fiber receives synaptic input from an inner hair cell via a single ribbon-type synapse with a single postsynaptic density, each expressing the Ca\(^{2+}\)-limiting AMPAR subunit GluA2 (Jing et al., 2013; Rutherford, 2015; Liberman and Liberman, 2016). Here, we show that these postsynaptic densities all coexpress GluA3 and GluA4 with GluA2 (Fig. 10). The total volume of AMPAR immunofluorescence was calculated for each synapse by thresholding a virtual channel created by summing the three individual confocal channels (GluA2, GluA3, and GluA4). Total AMPAR volume ranged from 0.09 to 0.85 μm\(^3\) per synapse. We calculated the intensity of each channel by summing the pixel values within each synaptic volume. Intensity of AMPAR sub-
Calcium-permeable AMPARs mediate synaptic currents at an adult auditory synapse

Our studies in fish and rats have demonstrated that CP-AMPARs mediate excitotoxic damage and may be expressed in mature mammalian hair cell synapses. However, to determine whether synthetically released glutamate can activate CP-AMPARs in vitro, paired recordings between single hair cells and their afferent fibers are necessary. However, this has proven to be extremely difficult in mature mammalian cochlea. We thus performed paired recordings between auditory hair cells and their afferent fibers in adult bullfrog amphibian papilla, an organ that detects airborne sounds of low frequency (100–1200 Hz; Smotherman and Narins, 2000; Li et al., 2014). Figure 12A shows an example recording of a hair cell and afferent fiber together with simultaneous time-resolved membrane capacitance measurements (a presynaptic measure of synaptic vesicle exocytosis; Li et al., 2009). The voltage-clamped hair cell was depolarized from −90 to −30 mV for 20 ms. This elicits a calcium current (ICa) and a Cm jump from the hair cell, and an EPSC in the afferent fiber (black trace). At this synapse, the EPSC charge is well correlated to the Cm jump (Cho et al., 2011). The EPSC is mediated by the opening of AMPA receptors in the afferent fiber (Keen and Hudspeth, 2006). It is composed of an initial phasic component followed by a tonic sustained component. This EPSC was partly blocked by 60 μM IEM, a use-dependent-specific open channel blocker of CP-AMPARs (Kim and von Gersdorff, 2016). Note that the large phasic peak of the EPSC was completely blocked after 7 min and 20 s in IEM, whereas the size of the Cm jump did not change (Fig. 12A).

The EPSC block by IEM was use dependent and progressive. Figure 12B shows the progressive block of the EPSCs in the continuous presence of IEM by successive 20 ms depolarizing pulses.

unit immunofluorescence per synapse scaled with volume over approximately one order of magnitude among synapses (Fig. 10B), suggesting differences among synapses in overall AMPAR abundance. Although each postsynaptic density expressed all three GluA subunits, we found heterogeneity suggesting that different types of AMPAR channels were present. Differences in fluorescence ratios generated the appearance of synapses with different hues in merged color panels (Figs. 10A, 11A, C). At some synapses, GluA2, GluA3, and GluA4 appeared to occupy very similar overlapping regions. Within other synapses, the subunits appeared to occupy partially nonoverlapping domains. Analysis with line profiles revealed some GluA2-lacking regions within synapses (Fig. 11B, arrowheads). To represent the subunit heterogeneity across synapses, we calculated fluorescence intensity ratios for each subunit compared with the sum intensities of all subunits at each synapse. While fluorescence ratios do not measure relative subunit abundance at a given synapse (see Materials and Methods), comparisons of these ratios among all synapses provide a measure of the heterogeneity of subunit relative abundance. Although the mean fluorescence ratios were similar for the population of synapses (0.31, 0.35, and 0.34, respectively, for GluA2, GluA3, and GluA4), the distributions of fluorescence ratios per synapse revealed that GluA2, GluA3, and GluA4 were expressed in different ratios at different postsynaptic densities (Fig. 11D). GluA4 exhibited the largest range of fluorescence ratios among synapses, from 0.23 to 0.47. In summary, the distribution of AMPAR subunits within and among synapses appears to be heterogeneous and is consistent with the presence of subregions with some GluA2 lacking CP-AMPARs, even though all synapses contain GluA2 subunits.
to the hair cell. The low affinity for glutamate of AMPARs, and perhaps the heterogeneous release probability of the synapse, may require repeated bouts of multiquantal glutamate release to produce complete block of EPSCs mediated by CP-AMPAR. Figure 12 shows that 60/100 M IEM reduced significantly the total charge of the EPSCs. After 5 min of IEM application, the average charge of EPSCs was decreased by 31.5 ± 4.2% (n = 16 pairs). Among these 16 pairs, 5 pairs showed smaller effects on the decrease of EPSC charges (13.0 ± 3.4%). After 8 min of IEM application, the average charge of the EPSCs was further decreased by 39.3 ± 6.0% (n = 8 pairs) without significant changes in the membrane capacitance jump. Because the large and phasic EPSC is responsible for rapid spike triggering in the afferent fibers (Schnée et al., 2013; Graydon et al., 2014), we also analyzed the amount of IEM block in the first 5 ms after the depolarizing pulse to the hair cell. During the first 5 ms, IEM decreased the EPSC charge by 44.9 ± 8.7%. These data clearly demonstrated that CP-AMPARs comprise a substantial component of the glutamatergic transmission at this mature auditory ribbon-type synapse.

Discussion

Using in vivo electrophysiology and Ca²⁺ imaging, we demonstrate that CP-AMPARs mediate glutamatergic transmission and excitotoxic damage at synapses between hair cells and their afferent neurons. Our study extends previous findings in acoustic-lateralis endorgans by demonstrating that CP-AMPARs mediate a major component of the afferent terminal synaptic response in adult auditory synapses of the frog. We demonstrate that excitotoxic damage of lateral line synapses in the zebrafish is associated with Ca²⁺ accumulation and that those synapses can be protected by pharmacological blockade of CP-AMPARs. Moreover, simultaneous immunohistochemistry for GluA2, GluA3, and GluA4 at mature afferent synapses in the mammalian cochlea suggests the existence of AMPA receptors lacking the Ca²⁺-limiting GluA2 subunit. Together, these findings suggest that CP-AMPARs are a common feature of primary afferent synapses in vertebrate acoustic-lateralis endorgans.
Presynaptic versus postsynaptic mechanisms of excitotoxic damage in the lateral line

Our findings with immunohistochemical (Fig. 8) and Ca\(^{2+}\)-imaging experiments (Fig. 4) suggest that the loss of terminal responsiveness following AMPA exposure is due to a loss of postsynaptic sensitivity instead of a restructuring of the synapse. Immunohistochemistry suggested that, despite substantial terminal swelling, the presynaptic and postsynaptic components remained intact at many synapses (Fig. 8). Yet it remains possible that the loss of postsynaptic terminal responsiveness following AMPA exposure was due to hair cell damage affecting presynaptic glutamate release (Sheets, 2017). Our current studies do not thoroughly address this possibility. However, lateral line hair cell responses to mechanical stimulation were intact after AMPA application (Fig. 5). This was assayed by water jet-evoked intracellular Ca\(^{2+}\) signals in hair cells, which are a measure of mechanotransduction and presumably a depolarized receptor potential (Kindt et al., 2012). The extent to which the loss of postsynaptic terminal responses resulted from the loss of presynaptic release from the stimulated hair cell, versus loss of postsynaptic sensitivity of the afferent fiber, remains to be explored.

Ca\(^{2+}\)-permeable AMPARs at auditory hair cell afferent synapses in the amphibian papilla

In addition to showing that CP-AMPARs contribute to AMPA-mediated Ca\(^{2+}\) influx in the fish, we have demonstrated that CP-AMPARs mediate a major component of glutamatergic transmission in auditory hair cell synapses of the bullfrog. Our finding that IEM1460 reduced the synaptic current charge by \~\(-40\%\) suggests that a large portion of the EPSC in the amphibian papilla is carried by CP-AMPARs (Fig. 12). In addition to Ca\(^{2+}\) permeability, the GluA2 subunit has been shown in other systems to influence several properties of AMPARs including EPSC kinetics (Geiger et al., 1995; Washburn et al., 1997). For example, AMPARs in the avian cochlear nucleus have high Ca\(^{2+}\) permeability and fast kinetics (Otis et al., 1995). EPSCs at hair cell afferent synapses have large amplitudes and submillisecond rise and decay kinetics (Glotzatki and Fuchs, 2002; Keen and Hudspeth, 2006), which are properties that promote spike phase locking to sound frequency (Li et al., 2014; Heil and Peterson, 2017). In addition to reducing the overall EPSC charge, IEM1460 greatly reduced and delayed the peak of the EPSC (Fig. 12A, inset). The initial phasic EPSC is thus mediated mostly by the opening of CP-AMPARs after synaptically released glutamate. As expected for a use-dependent open channel blocker of CP-AMPARs, this reduction by IEM1460 was activity dependent (Fig. 12B). It remains to be seen whether CP-AMPARs carry a significant fraction of the EPSC in terminals of auditory nerve fibers in the mammalian cochlea, which are postsynaptic to inner hair cell synaptic ribbons.

GluA2-lacking AMPARs at hair cell synapses in the mammalian cochlea

Cochlear hair cell afferent synapses contain post synaptic AMPAR subunits including the Ca\(^{2+}\)-limiting GluA2 subunit (Fig. 11). Expression of the GluA2 subunit juxtaposed to the presynaptic ribbon is used to define a synapse, and count them, over development and in response to acoustic trauma (Liberman et al., 2015; Liberman and Liberman, 2016). The presence of GluA2 at all synapses has contributed to the general assumption that AMPARs in the mammalian cochlea are relatively impermeable to Ca\(^{2+}\).

AMPA-stimulated Ca\(^{2+}\) entry into postsynaptic terminals has been proposed as the cause of excitotoxic damage to auditory nerve fibers, primarily because Ca\(^{2+}\) influx through glutamate receptors is a well studied excitotoxic mechanism in the brain (Sattler and Tymianski, 2001). However, its relevance in the inner ear is unclear. Cobalt staining, used as a proxy for Ca\(^{2+}\) entry (Pruss et al., 1991), has suggested that AMPA induces Ca\(^{2+}\) entry into SGNs (Eybalin et al., 2004). However, given that kainate receptors, NMDA receptors, and voltage-gated Ca\(^{2+}\) channels may mediate Ca\(^{2+}\) influx as well, it is still unclear whether CP-AMPARs are directly involved in cochlear excitotoxicity.

Acoustic overstimulation results in swelling of auditory nerve fiber terminals on hair cells, which express GluA2, GluA3, and GluA4 (Fig. 10). With confocal microscopy, we found GluA2-lacking regions within postsynaptic densities of some auditory nerve fiber terminals in the rat (Fig. 11). This suggests that, although all postsynaptic densities expressed GluA-2, some of the AMPAR tetrmers may lack the GluA2 subunit. On most postsynaptic densities, the three AMPAR subunits appeared to overlap each other at confocal resolution. Future work with super-resolution light microscopy is needed to assess the prevalence of GluA2-lacking regions within synapses. At central synapses including those in the calyx of Held in the mouse auditory brainstem, the presence of GluA4 in GluA3/4-containing AMPARs is indispensable for driving fast neu-
rotransmission (Yang et al., 2011). The existence of functional GluA2-lacking CP-AMPARs and their role in cochlear synaptic transmission remains to be tested. At hair cell synapses, relatively fast kinetics and Ca2+-permeability of GluA2-lacking AMPARs may be important for fast auditory signaling during phase locking as well as for understanding mechanisms of excitotoxicity upon overexposure to sound.

References
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