Excessive activation of ionotropic glutamate receptors induces apoptotic hair-cell death independent of afferent and efferent innervation

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Accumulation of excess glutamate plays a central role in eliciting the pathological events that follow intensely loud noise exposures and ischemia-reperfusion injury. Glutamate excitotoxicity has been characterized in cochlear nerve terminals, but much less is known about whether excess glutamate signaling also contributes to pathological changes in sensory hair cells. I therefore examined whether glutamate excitotoxicity damages hair cells in zebrafish larvae exposed to drugs that mimic excitotoxic trauma. Exposure to ionotropic glutamate receptor (iGluR) agonists, kainic acid (KA) or N-methyl-D-aspartate (NMDA), contributed to significant, progressive hair cell loss in zebrafish lateral-line organs. To examine whether hair-cell loss was a secondary effect of excitotoxic damage to innervating neurons, I exposed neurog1a morphants—fish whose hair-cell organs are devoid of afferent and efferent innervation—to KA or NMDA. Significant, dose-dependent hair-cell loss occurred in neurog1a morphants exposed to either agonist, and the loss was comparable to wild-type siblings. A survey of iGluR gene expression revealed AMPA-, Kainate-, and NMDA-type subunits are expressed in zebrafish hair cells. Finally, hair cells exposed to KA or NMDA appear to undergo apoptotic cell death. Cumulatively, these data reveal that excess glutamate signaling through iGluRs induces hair-cell death independent of damage to postsynaptic terminals.

Intense acoustic trauma or ischemic injury leads to accumulation of the excitatory neurotransmitter glutamate in the cochlea1-4. There is evidence that excess glutamate acts as a primary trigger for subsequent pathologies in noise-exposed cochlea, the most well-characterized effect being consequent swelling of postsynaptic afferent nerve terminals resulting from overactivation of AMPA-type GluRs5-8. By contrast, whether excess glutamate signaling damages hair cells—the sensory receptors of the auditory system—has not yet been fully examined. Presynaptic iGluRs that regulate neurotransmitter release have been observed in many areas of the central nervous system9, and several studies suggest that all three types of iGluR subunits—AMPA, Kainate, and NMDA—are expressed and presynaptically-localized in cochlear hair cells10-14. Yet whether excessive activation of iGluRs contributes to hair-cell damage has not been directly studied in a mammalian model system because it is difficult to discern whether hair-cell death in iGluR-agonist exposed cochlea is the result of damage to the hair cells themselves or collateral damage from injured postsynaptic nerve terminals15.

Zebrafish afford a useful model system to address whether glutamate toxicity damages sensory hair cells. Zebrafish hair cells are homologous to mammalian hair cells16-20, yet are optically accessible in whole larvae within the lateral line organ—a sensory organ used to detect the movement of water that contains clusters of superficially localized hair cells called neuromasts (NMs). Additionally, zebrafish hair cells are amenable to pharmacological manipulation, allowing for drug application and subsequent examination of hair-cell morphology and function. This is particularly advantageous for investigating hair-cell toxicity, as delivering drugs into the cochlea is challenging and can in and of itself damage sensory hair cells21.

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I therefore determined whether glutamate excitotoxicity directly damages hair cells by examining lateral-line NMs of 5 to 6-day-old zebrafish larvae exposed to drugs that mimic glutamate-induced excitotoxic trauma. Exposure to the iGluR agonists kainic acid (KA) or N-methyl-D-aspartate (NMDA) contributed to significant, progressive hair-cell loss in both wild-type larvae and in neurogenin1 morphants—fish that have morphologically mature hair cells devoid of afferent and efferent innervation. Analysis of iGluR expression in isolated hair cells populations subsequently revealed that, similar to what has been previously reported in mammalian systems, AMPA-, Kainate and NMDA-type receptor subunits are expressed in zebrafish hair cells. KA and NMDA mediated hair-cell death is characterized by the formation of apoptotic bodies and activation of caspase-3. Cumulatively, these data indicate that excessive signaling through iGluRs induces apoptotic hair-cell death, and suggests cell death may be instigated through iGluRs on the hair cells themselves.

Results

KA exposure leads to swelling and bursting of postsynaptic afferent terminals. There is an abundance of evidence that cochlear nerve fibers are damaged by exposure to iGluR agonists: previous studies have reported excitotoxic damage to cochlear nerve fibers akin to that brought about by noise overexposure in cochlea briefly treated with the agonist α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA)2,22 or the more potent excitotoxic agonist KA23–25. To confirm whether zebrafish lateral-line afferent neurons are similarly sensitive to AMPA/KA GluR agonist-induced excitotoxic trauma, I exposed, to KA, live 6-day-old transgenic zebrafish larvae expressing GFP in their afferent neurons26 and mcherry at the hair-cell presynaptic ribbons while recording changes in their afferent terminal morphology using confocal time-lapse imaging. I observed profound swelling of lateral-line afferent terminals (Fig. 1A; white arrowheads) analogous to that observed in KA exposed mammalian cochlea23,27. In addition, I applied the GluR agonist NMDA to these transgenic larvae and did not observe swelling of afferent terminals (Fig. 1B), which is also consistent with what had been previously observed in drug-perfused mammalian cochleae. These observations support that iGluR agonists act on lateral-line afferent neurons in a similar manner as auditory dendrites.

KA or NMDA exposure initiates NM hair-cell loss. Subsequently, I examined lateral-line NM hair-cell morphology in 5 to 6-day-old larvae that were exposed to AMPA, KA, or NMDA, then either immediately fixed or rinsed and allowed to recover either 2 hours or overnight prior to fixation. Unexpectedly, exposure to these high (300μM) levels of either KA or NMDA contributed to significant, progressive hair cell loss (Fig. 1C–F) i.e. exposure to either KA or NMDA lead to noticeable hair-cell loss (Fig. 1E) that became more significant two hours after drug exposure (Fig. 1F). This result was surprising—the presence of functional iGluRs in sensory hair cells has been inferred by previous studies but not definitively established, so iGluR overactivation has not been considered as a cause of hair-cell death. Considering that KA exposure leads to swelling and bursting of afferent synaptic contacts (Fig. 1A), I also examined whether exposure to KA or NMDA lead to hair-cell synapse loss, but did not find any significant difference in the number of presynaptic ribbons per remaining NM hair cell in KA or NMDA exposed larvae (Fig. 1H). Cumulatively, these data show that activation of AMPA/Kainate or NMDA type receptors in zebrafish lateral line organs contributes to similar neural pathological changes as that observed in mammalian cochlea, and support that overactivation of iGluRs, possibly on the hair cells themselves, also contributes to noise-induced hair-cell damage and death.

neurogenin1 morphant NM hair cells appear morphologically mature and have functional mechanotransduction. To exclude the possibility that damage to hair cells from exposure to KA or NMDA was an indirect result of overstimulating lateral-line nerve fibers, I examined neurogenin1 (ngn1) morphants: larvae whose hair-cell organs are devoid of afferent and efferent innervation28 (Fig. 2B). Zebrafish lacking ngn1 are similar to mouse ngn1 mutants in that their hair-cell organs lack innervation29,30. Ngn1-lacking zebrafish survive past the onset of lateral-line hair-cell function30 due to nutrition provided by the yolk sack. Prior to exposing the larvae to drugs, I examined whether, in 5-day-old ngn1 morphants, NM hair cells had the morphological characteristics of functionally mature hair cells. Immunolabeled ngn1 morphant hair cells showed several known hallmarks of morphological maturity: they had basally-localized presynaptic ribbons that localized with the synaptic-vesicle marker Vglu3 (Fig. 2D) as well as clusters of L-type calcium channels Cav1.3 localized at the synaptic-ribbon (Fig. 2E). I also examined functional maturity by briefly exposing ngn1 morphants to the vital dye FM1–43 at a concentration that has been previously shown to preferentially label relatively mature lateral-line NM hair cells with mechanotransduction activity31. Brief exposure to low concentrations of FM1–43 is a robust assay for functional hair-cell mechanotransduction that has been verified in both mammalian and zebrafish systems32,33. I observed that ngn1 morphant NM hair cells take up FM1–43 (Fig. 2G), supporting that ngn1 morphant zebrafish hair cells are morphologically and, at least in part, functionally mature despite lacking afferent and efferent innervation. It should be noted that hair-cell innervation is important for synaptic-rhombomere maintenance in zebrafish34 and afferent innervation plays a significant and variable role in long-term hair-cell maintenance and viability in mice35. Nevertheless, previous studies have shown that short term hair-cell survival is possible in the absence of innervation36, indicating ngn1 morphants as a useful tool in determining whether iGluR agonist exposure damages hair cells in the absence of innervation.

KA or NMDA exposed ngn1 morphants show NM hair-cell loss that is comparable to wild-type and dose dependent. Consequently, I exposed 5-day-old ngn1 morphants and their WT siblings to KA or NMDA for 1 h., then allowed them to recover for 2 h. prior to fixation. Significant hair-cell loss occurred in ngn1 morphants exposed to 300μM KA or NMDA (Fig. 3D–H), and the loss was comparable to that observed in wild-type siblings (Fig. 3A–C,G). The actions of KA and NMDA are likely specific, as hair-cell loss in ngn1 morphants occurred in a dose-dependent manner (Fig. 4A,B). Additionally, KA-induced hair-cell loss was blocked...
Figure 1. Exposure to KA or NMDA initiates hair-cell loss, but not synaptic-ribbon loss. (A,B) Live-imaging of KA (A) or NMDA (B) exposure: Representative frames from a movie taken over 2 hours with one frame per 3.5 minutes. Each frame is a maximum intensity projection (z-stack top-down image) of posterior lateral-line (LL) NM3 in a 6 dpf transgenic larva expressing Ribeye a-tagRFP in hair cells and GFP inafferent nerve fibers. Arrowheads in (A) indicate afferent-terminal swellings; asterisks indicate bursting afferent terminals. (C,D) Representative maximum intensity top-down (x-y) projections of immunolabeled Vglut3 (red) labeling the basolateral-end of hair cells and Ribeye b (green) labeling synaptic ribbons in NM3 of 6 dpf larvae. Larvae were exposed to DMSO carrier alone (Control) or iGluR agonists for 1 hour, then either immediately fixed ((C); No rec.) or rinsed and allowed to recover for 2 hours ((D); 2h. rec.). Hair-cell death was confirmed by absence of DAPI-labeled nuclei. Scale bars: 3 μm (E–H) The number of intact hair cells per NM (E–G) or number of synaptic ribbons per hair cell (H) in 6 dpf (E,F,H) or 7 dpf (G) larvae exposed to DMSO alone or 300 μM iGluR agonist. Each circle represents NM3 in an individual larva. There was no significant loss of hair cells in NMs following exposure to AMPA at any time-point or the more excitotoxic AMPA/Kainate receptor agonist KA immediately following exposure (E). However, there were significantly fewer hair cells per NM in KA and NMDA-treated larvae compared to control 2 hours after drug exposure (F; ***p < 0.0001 defined by Dunnett’s multiple comparisons test) and after 20–22 h. recovery post-exposure (G; **p < 0.01 defined by unpaired t test). There was no significant difference in the number of synaptic ribbons per hair cell 2 h. after drug exposure (H), defined by Dunnett’s multiple comparisons test).

when morphants were co-exposed to the competitive AMPA/kainate receptor antagonist CNQX (Fig. 4A) and NMDA-induced hair-cell loss was partially blocked by NMDA receptor antagonist D-AP5 (Fig. 4C). The actions of KA and NMDA appeared to be synergistic; a non-lethal concentration of 30 μM KA combined with a minimally lethal concentration of 30 μM NMDA lead to significantly more hair-cell death than 30 μM NMDA alone (Fig. 4D; Control = 11 ± 0.4, 30 μM KA = 10 ± 0.6, 30 μM NMDA = 8 ± 0.3; 30 μM KA + 30 μM NMDA = 5 ± 0.4). To analyze the synergistic effect of KA and NMDA combined, I calculated the coefficient of drug interaction (CDI; see Methods)39. The CDI of KA + NMDA (30 μM) was 0.7; CDI < 1 indicates synergism. Cumulatively, these data
suggest that excessive glutamate signaling mediates damage to sensory hair cells independent of damage to post-
synaptic terminals, and implies the presence of functional iGluRs on hair cells.

Zebrafish hair cells express AMPA-, KA-, and NMDA-type receptor subunits. Data from previous
studies suggest that mammalian hair cells express AMPA,10,11 Kainate-12, and NMDA-type14,40 GluR subunits.
To examine whether zebrafish hair cells express iGluR subunits, I performed gene expression analysis on hair
cells isolated from 5-day-old stable transgenic larvae. Fluorescence-activated cell sorting (FACS) of lysed lar-
val populations was used to separate mcherry containing hair cells and GFP containing neurons from other
non-fluorescent cell types (Fig. 5A). To perform a survey of iGluR expression, cDNA templates were created
from the sorted cells via RT-PCR, then used to generate small amplicons (~150–300 bp) of GluR transcripts of
the AMPA, Kainate, and NMDA receptor classes as well as cell-type specific control genes. Using this strategy, I
identified several iGluR subunits in zebrafish hair cells: AMPA-receptor subunit gria4b, Kainate-receptor subu-
nits grik1a and grik4, and NMDA-receptor subunits grin1a and grin2ba (Fig. 5B).

In addition, I detected immunolabeling of three identified iGluR subunits at hair-cell ribbon synapses. The
commercial antibody used to detect zebrafish Grik1a was generated using a 49aa region of the C-terminus of
human Grik2, which has 92% sequence conservation with zebrafish Grik1a aa 652–701 (ClustalW Method). I
observed Grik1a (Fig. 5C), Grik4 (Fig. 5D), and Grin1a (Fig. 5E) immunolabeled puncta localized to hair-cell
ribbon synapses. To determine the labeling distribution of iGluRs relative to both pre- and post-synaptic com-
ponents, I compared iGluR labeling intensity plotted in relation to distance (profile plots) with that of synaptic
ribbons and the afferent postsynaptic densities (PSDs). Figure 5(C–E) shows high-power images of representative
ribbon synapses and their corresponding profile plots. Kainate-receptor subunits Grik1a (Fig. 5C) and Grik 4
(Fig. 5D) labeling showed multiple peak intensities that overlapped with both Ribeye and PSD peak intensities,
suggesting these iGluR subunits localize to both pre- and postsynaptic regions. By contrast, NMDA-receptor

Figure 2. Characterization of ngn1 morphant NM hair cells. (A,B) Representative max intensity top-down
(x-y) and side-view (x-z) projections of imunolabeled Parvalbumin (red) labeling hair cells, acetylated-Tubulin
(green) labeling neurons and the apical-region of hair cells, and DAPI labeling nuclei in posterior LL NM2 of
WT (A) or ngn1 MO (B) 5 dpf zebrafish larvae. Note the absence of immunolabeled green neurites beneath
hair cells in the ngn1 morphant NM (x-z) image. Scale bar: 3 μm (C,D) Representative immunoabeled (x-y)
and (x-z) cross-sections of a WT (A) or ngn1 MO(A) anterior NM. Both synaptic ribbons (Ribeye; red) and
synaptic vesicles (Vglut3; white) localize to the basolateral end of ngn1 MO hair cells, despite the lack of afferent
innervation (HNK-1; green). Scale bar: 3 μm (E) Max-intensity(x-y) projection of immunolabeled synaptic
ribbons (red) and the presynaptic calcium channel Ca,v,1.3a (green) in NM3 of a 5 dpf ngn1 morphant larva.
Insets show colocalization of Ribeye and Ca,v,1.3, which is a hallmark of hair-cell maturation. Scale bars: 3 μm,
1 μm (insets) (F,G) Representative lateral views of 5 dpf WT (F) or ngn1 morphant (G) larvae briefly exposed
FM1-43. Insets show higher magnification top-down views of the NMs indicated by the white dashed squares.
Labeling with FM1-43 indicates functional mechanotransduction machinery in ngn1 morphant hair cells.
subunit Grin1 showed a single peak intensity that overlapped with Ribeye labeling the presynapse (Fig. 5E). To determine whether iGluR subunit labeling persisted in the absence of the postsynapse, I examined iGluR subunits labeling in ngn1 morphant fish, and observed Grik1a, Grik 4, and Grin1a labeling localized with Ribeye (Fig. 5F–H). These results indicate iGluR subunits are expressed in zebrafish hair cells, and suggest that KA and NMDA damage to hair cells may be mediated by hair cell autoreceptors.

KA or NMDA-induced hair-cell death is not accompanied by intracellular calcium ([Ca$^{2+}$]$_{i}$) dysregulation. One mechanism of excitotoxic cell death is dysregulation of [Ca$^{2+}$]$_{i}$ homeostasis followed by the loss of mitochondrial membrane potential$^{41}$. To examine whether [Ca$^{2+}$]$_{i}$ dysregulation occurs in hair cells exposed to KA or NMDA, I exposed ngn1 morphant larvae stably expressing the genetically encoded calcium indicator GCaMP3 in hair cells$^{42}$ to either KA or NMDA while recording changes in GCaMP3 fluorescence intensity using confocal time-lapse imaging. To verify I could visualize [Ca$^{2+}$]$_{i}$ dysregulation in ngn1 morphant hair cells, I exposed them to 50 μM gentamycin, which has been previously shown in this transgenic line to induce a marked increase in cytosolic [Ca$^{2+}$]$_{i}$ prior to hair-cell death$^{43}$. Likewise, I also observed sharp peaks in GCaMP3 fluorescence relative to baseline in gentamycin exposed hair cells prior to their death and extrusion from the NM (Fig. 6A,B). By contrast, I observed very little change in GCaMP3 fluorescence relative to baseline in dying hair cells exposed to either KA (Fig. 6C,D) or NMDA (Fig. 6E,F); dying hair cells exhibited small increases in GCaMP3 fluorescence prior to cell death (Fig. D (Cell 3); Fig. F (Cell 2)), but they were ~9 fold smaller than the calcium peaks observed in dying gentamycin-treated hair cells (Fig. 6B).

Figure 3. KA or NMDA exposure initiates hair-cell loss in ngn11 morphants comparable to WT siblings. (A–F) Representative max intensity top-down (x–y) and side-view (x–z) projections of immunolabeled Parvalbumin (red), acetylated Tubulin (green), and DAPI (blue) in posterior LL NM7 of 5 dpf WT (A–C) or ngn1 MO (D–F) larvae. Larvae were exposed to DMSO carrier alone (Control) or iGluR agonists for 1 hour, then rinsed and allowed to recover for 2 hours. Exposure to 300 μM KA (B,E) or 300μM NMDA (C,F) leads to hair-cell loss in both WT and morphants. White arrowheads indicate pyknotic nuclei. Scale bar: 3 μm (G,H) The number of intact hair cells per NM in 5 dpf larvae exposed to DMSO alone, 300 μM KA, or 300 μM NMDA. Each circle represents NM2 or NM7 in an individual larva. There were significantly fewer HCs per NM in KA and NMDA-treated larvae compared to control 2 h. after exposure in both WT siblings (G) and ngn1 (H) MO larvae (*p < 0.05; ***p < 0.001; ****p < 0.0001 defined by Dunnett’s multiple comparisons test).
The majority (>90%) of total Ca\(^{2+}\) current in hair cells is carried by the L-type calcium channel CaV1.3.\(^{18,31,43,44}\) To further examine whether [Ca\(^{2+}\)]\(_i\) dysregulation contributes to iGluR mediated hair-cell death, I inhibited extracellular calcium influx with the L-type calcium channel blocker isradipine at a concentration (10 μM) that has been previously shown to reduce calcium responses in zebrafish NM hair cells.\(^{45}\) Consequently, I observed a dramatic reduction in GCaMP3 fluorescence in NM hair cells co-exposed to isradipine (Fig. G,H). Notably, blocking extracellular calcium influx did not provide a protective effect; the NM co-exposed to NMDA and

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**Figure 4.** KA and NMDA induce hair-cell loss in a dose-dependent manner and have a synergistic effect in ngn1 morphants. (A,B) Dose-dependent hair-cell loss in neurog1 MO larvae exposed to KA (A) or NMDA (B). Each dot represents the mean hair cell loss per NM in 5–10 individual larvae; the bars represent the s.e.m. Co-exposure with the AMPA/KA receptor antagonist CNQX (300μM) blocks KA-induced hair-cell loss (A). (C) NMDA-exposed NM hair cells are partially protected from NMDA mediated death when pre-exposed to the competitive NMDA receptor antagonist D-AP5. Each circle represents NM2 or NM7 in an individual larva. (**p < 0.01; ****p < 0.0001 defined by Dunnett's multiple comparisons test). (D) The effect of combined KA and NMDA exposure on NM hair-cell death was greater than the sum of the effects of each drug alone. There were significantly fewer HCs per NM in larvae treated with a sub-lethal concentration of KA (A) and NMDA than NMDA alone. (**p < 0.01; ****p < 0.0001 defined by Dunnett's multiple comparisons test).
isradipine lost hair cells (Fig. G; white dashed outline), which was never observed in NM hair cells exposed to isradipine alone. Cumulatively, these data support that KA and NMDA do not bring about hair-cell death through dysregulation of $[\text{Ca}^{2+}]_I$ homeostasis and mitochondrial collapse.

**KA or NMDA exposed ngn1 morphant NM undergo apoptotic cell death.** Hair cells exposed to acoustic trauma have been shown to undergo apoptotic cell death, necrotic cell death, or a combination of the two. To determine whether hair-cell death in iGluR-agonist exposed hair cells appears characteristically apoptotic or necrotic, I examined changes in hair-cell morphology in ngn1 morphant larvae to either KA or NMDA. Exposure to either KA (Fig. 6C; Supplemental Fig. 1A) or NMDA (Fig. 6E; Supplemental Fig. 1B) led to the formation of blebs and what appeared to be apoptotic bodies accompanied by hair-cell extrusion from the NMs. I subsequently examined whether caspase-3—a downstream effector of apoptotic cell death—was activated in hair cells exposed to high concentrations (300 μM) of KA or NMDA, and found the majority of NMs contained a subset of hair cells with activated caspase-3 immediately following drug exposure (Fig. 7A–D). Additionally, I
examined supporting cell morphology and caspase-3 labeling in the transgenic fish line Supporting-Cell Marker 1 (scm1:GFP), which expresses GFP in all the supporting cells of the lateral line47. NM supporting cells appeared intact and their gross morphology comparable to unexposed control immediately following iGluR agonist exposure (Fig. 7E–G). Activated caspase-3 labeling corresponded with hair-cell labeling (Fig. 7F, G), suggesting that activation of downstream cell death pathways in hair cells is not indirectly mediated by supporting-cell damage or death. These data reveal that ngn1 morphant NM hair cells exposed to KA or NMDA undergo caspase 3-mediated cell death that appears apoptotic.

Discussion
It has been well established that glutamate excitotoxicity resulting from ischemia or acoustic trauma contributes to cochlear nerve-terminal damage. My observations in the zebrafish lateral-line support the idea that glutamate excitotoxicity also contributes to hair-cell damage and death. The significant findings of this study are i) overactivation of AMPA/Kainate or NMDA-type GluRs leads to hair cell loss, ii) hair-cell loss is independent of afferent...
and efferent innervation, iii) zebrafish NM hair cells express AMPA/Kainate and NMDA-type GluR subunits and, iv) overactivation of iGluRs contributes to apoptotic hair-cell death. These observations in the zebrafish lateral line may give insight into the mechanisms of cochlear ischemic injury, which can result from loud noise exposure. Previous studies support that transient cochlear ischemia promotes hair-cell loss, particularly the loss of inner hair cells. It has been proposed that delayed inner-hair cell death following cochlear ischemia is due to excessive glutamate accumulation; glutamate levels in the perilymph increase considerably following ischemic injury, perfusion of the AMPA/Kainate receptor antagonist DNQX protects against inner hair cell loss following transient cochlear ischemia, and treatment of cochlear explants with NMDA receptor blockers protects against hypoxia-induced hair-cell loss. Additionally, inner hair cells have been shown to undergo apoptotic cell death following transient cochlear ischemia, which corresponds to what I observed in iGluR agonist treated zebrafish lateral line. Speculatively, these previous reports combined with the results of this study suggest a model whereby cochlear ischemia and subsequent excess glutamate accumulation may result in over-activation of both AMPA/Kainate- and NMDA-type receptors initiating apoptotic hair-cell death.

The presence of iGluRs in hair cells suggests that activation of autoreceptors on the hair cells themselves initiates intracellular signaling cascades that lead to hair-cell apoptosis. The classical pathway by which glutamate toxicity initiates cell death is by excessive calcium influx leading to mitochondrial dysfunction and the generation of caspase-3. The following figure (Figure 7) shows representative images of NM7 in 5 dpf ngn1 morphant larvae exposed to DMSO alone, 300 μM KA, or 300 μM NMDA, then immediately processed for histology. NM7 exposed to iGluR agonists stained positively for activated Caspase-3, indicating the cells were undergoing apoptotic cell death. A floating bar graph showing the percentage of NM in individual larval posterior lateral line that stained positive for activated Caspase-3. Each bar represents the min. and max. values in 11–15 larvae; the lines represent the mean values. (****p < 0.0001 defined by Dunnett’s multiple comparisons test).
of free radicals. Notably, I observed very little change in \([\text{Ca}^2+]_i\) in dying hair cells exposed to KA (Fig. 6C,D) or NMDA (Fig. 6E,F) and no protective effect from blocking extracellular calcium influx through \(\text{Ca}_\text{v}1.3\) channels (Fig. 6G,H), indicating that dysregulation of intracellular calcium is not what drives hair-cell death following excess iGluR activation. There is substantial evidence that specific iGluRs are coupled to the activation of distinct signaling pathways, and it is the triggering of these pathways though excess activation of the receptor rather than an overall elevation in intracellular calcium levels that leads to the activation of programmed cell death. This is further supported by my observations that KA and NMDA have a synergistic effect on hair-cell loss (Fig. 4D), indicating that KA and NMDA initiate hair-cell death through distinct pathways. Additional studies will be needed to verify that hair-cell localized iGluRs conduct calcium current, and to identify the specific downstream signaling pathways they trigger when overactivated.

What might be the physiological role of presynaptic iGluRs in sensory hair cells? Based on previous observations in the central nervous system, presynaptic iGluRs may modulate hair-cell synaptic transmission in response to activity. Presynaptic iGluRs examined in the brain have been shown to generally facilitate synaptic transmission. For example, kainate autoreceptors localized to hippocampal mossy fiber presynaptic active zones facilitate subsequent release of glutamate when activated, and Grin2B containing NMDA autoreceptors in the entorhinal cortex tonically facilitate glutamate release during development. On the other hand, presynaptic iGluRs have also been shown to depress evoked transmitter release; activation of kainate receptors in dorsal root ganglion neurons suppresses their glutamate release. It will be interesting to determine in future studies whether presynaptic iGluRs modulate glutamate release from sensory hair cells and, if so, in what way they affect synaptic transmission.

In conclusion, sensory hair cells are vulnerable to glutamate excitotoxicity and may be damaged via excess activation of presynaptic iGluRs. The results of this study reveal a potential mechanism of hair-cell death from excitotoxic cochlear injury. Excess glutamate accumulation and excitotoxicity are not unique to ears; glutamate excitotoxicity is responsible for the activation of diverse cell-death pathways in hypoxic-ischemic brain injury as well as a number of neurodegenerative diseases. Determining the mechanisms of excitotoxic pre- and postsynaptic damage in hair-cell organs may provide a better understanding of excitotoxic pathways underlyng a variety of neuropathological conditions.

Methods

Ethics Statement. This study was performed with the approval of the Massachusetts Eye and Ear Animal Care Committee and in accordance with NIH guidelines for use of zebrafish.

Fish Strains and Husbandry. Both larval and adult zebrafish were maintained in 14-hour light, 10 hour dark cycle. Adult fish were fed twice daily with GEMMA Micro 300 (Skretting). Embryos for experiments were obtained through paired-mating of Tübingen wild-type fish or the following transgenic lines: neurod:GFP, -\(\text{ngn1}\); ribeyea-mcherry, -\(\text{myosin6b:ribeyea-mcherry}\), -\(\text{myosin6b:gcamp}\), and -\(\text{scm1:GFP}\) and were maintained in E3 embryo media (5 mM NaCl, 0.17 KCl, 0.33 mM CaCl2 and 0.33 mM MgSO4) at 29°C.

Statistical analysis. Statistical analysis was performed using Prism 5 (GraphPad Software Inc). The Kolmogorov-Smirnov test was used to test and confirm the normality of data distributions. Statistical significance between multiple conditions was determined by one-way ANOVA and the appropriate post-hoc test; statistical significance between two conditions was determined by unpaired Student’s t tests.

Primary Antibodies. The following commercial antibodies were used in this study: MAGUK (K28/86; 1:500; NeuroMab, UC Davis), acetylated Tubulin (1:1000; Sigma-Aldrich), Parvalbumin (1:2000; Thermo Scientific), HNK-1 (Zn12; 1:500; DSHB, University of Iowa), Otoferlin (HCS-1; 1:500; DSHB, University of Iowa), cleaved Caspase-3 (1:400; Cell Signaling Technology), Gria 4 (1:1000; Synaptic Systems). In addition, the following affinity-purified antibodies generated against Danio rerio (International), Grik 4 (1:400; Genway Biotech Inc.), Grin 1 (1:1000; Synaptic Systems). In addition, the following affinity-purified antibodies generated against Danio rerio were also used in this study: Ribeye b (mouse IgG2a; 1:2000), Vglut3 (rabbit polyclonal; 1:1000), and Ca\(_\text{v}1.3\)a (rabbit polyclonal; 1:1000).

Whole-Mount Immunohistochemistry and Fluorescence Imaging. 5–6 day old larvae were larvae were quickly sedated on ice, then transferred to fixative (4% paraformaldehyde/4% sucrose/0.15 mM CaCl2 in phosphate buffer) for either 1 hour (Gria and Grik immunolabel) or 6 hours at 4°C. Larvae immunolabeled with antibodies for Gria or Grik subunits were permeabilized in 1% Tween in phosphate buffered saline (PBS) 5 hours at 4°C, then blocked in PBS buffer containing 1% bovine serum albumin (BSA), 0.5% Fish Skin Gelatin, 2% goat serum, and 0.1% dimethyl sulfoxide (DMSO). All other larvae were permeabilized with ice cold acetone for 5 minutes, then blocked with PBS buffer containing 2% goat serum, 1% BSA, and 1% DMSO. All immunos were incubated with primary antibodies diluted in blocking buffers overnight followed by diluted secondary antibodies coupled to Alexa 488, Alexa 647 (Molecular Probes, Invitrogen), or DyLight 549 (Jackson ImmunoResearch), and labeled with DAPI (Molecular Probes, Invitrogen). Z-stack images of NMs (spaced by 0.3 μm) were acquired with a Leica SP8 confocal microscope using a 63X 1.3 N.A. 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 (FEI).

Morpholino injection. neurogenin 1 morphant larvae have previously been shown to have a large loss of cranial ganglia as neurogenin1 mutants. MOs against the start codon of ngn1 (5’-CCATATCGGAGTAGTAC GATCTCCAT-3’) were obtained from GeneTools (Philomath, OR) and used as previously described. Approximately 1–2 nl of 0.84 mM ngn1 MO diluted in RNAse-free double distilled water with 3% phenol red was pressure injected.
into 1-cell stage wild-type and Tg(neurod:GFP) outcross embryos. Loss of lateral-line ganglia was confirmed by visualizing the absence of either GFP-labeled neurites in neurod:GFP transgenics or acetylated tubulin immunolabel.

**Pharmacological manipulation of larvae.** All of the drugs used in this study were acquired from Abcam. 5–6 days post fertilization (dpf) larvae were exposed to drugs diluted in E3 with 0.1% DMSO for 50 minutes at 29°C for histology or at room temperature (20–22°C) during live imaging. E3 with 0.1% DMSO were used as controls. Following drug exposure, larvae were either immediately fixed for immunohistochemistry or allowed to recover for 2 hours or overnight at 29°C. To determine if there was an additive or synergistic effect between KA and NMDA, the coefficient of drug interaction was calculated per11 using the following equation: CDI = AB/(A × B) whereby AB is the ratio of the combined drug group to the control group, and A or B is the ratio of the single drug group to the control group. CDI < 1 indicates synergism, CDI = 1 indicates additivity, and CDI > 1 indicates antagonism. For 30μM KA + 30μM NMDA: ((5/11)/(10/11)/(8/11)) = 0.45/0.66 = 0.68 (~0.7).

**FM1-43 Labeling.** Free-swimming 5 dpf larvae were immersed in E3 medium containing 3μM FM 1–43 (n-(3-triethylammoniumpropionyl)-4-(4-(dibutylamino)-styryl) pyridinium dibromide; Molecular Probes, Invitrogen) for 30 s, followed by three rinses in E3 (as described in ref. 32). Larvae were then anesthetized with 0.2 mg/ml of tricaine methanesulfonate (Sigma-Aldrich), placed in a depression slide, and imaged with a Nikon E800 Microscope using 10x/0.45 NA and 40x/0.95 NA dry objective lenses.

**Live-imaging of afferent neurons.** 5–6 dpf larvae were anesthetized with 0.2 mg/ml of tricaine, then soaked in 1 mg/ml α-bungarotoxin (Alamone labs) for 2 minutes and rinsed in E3. They were kept in E3 prior to imaging until they no longer swim or responded to touch (~20 min.). Larvae were mounted lateral-side on a thin layer of 1–2% low-melt agarose in a 35 mm dish and covered in E3 media. Z-stack images (spaced by 0.6 μm over 5–10 μm) were acquired with a Leica SP2 confocal microscope using a 60x/0.9 N.A. water immersion lens.

**Calcium imaging.** 5 dpf ngn1 MO larvae were anesthetized and maintained in 0.2 mg/ml of tricaine or soaked for 5 seconds in 0.6 mg/ml pancuronium bromide (Sigma-Aldrich) and maintained in E3 solution. They were immobilized for imaging in a quick release open bath chamber with a brain-slice harp (Warner Instruments). Methods for imaging and analyzing myo6b:gcamp3 larvae were based on those previously described using this transgenic line47. Images were taken with a Leica SP8 equipped with a HyD hybrid detector at 1400 Hz/6 fps. A 488 Argon laser was used for GCAMP excitation; the laser power was adjusted such that pixel intensities were set at <25% saturation. Baseline readings were taken at 30 sec intervals for 5 min. For drug application, 1/4th of the E3 bath solution was removed and carefully replaced with 4X concentrated drug solution. Additional readings were taken every 30 sec for 50 minutes during drug application. 4-D stacks were converted into maximum intensity projections in ImageJ. Lateral movements (X-Y) of the NMs were corrected using the StackReg plugin (as described in ref. 69). Following image registration, temporal changes in fluorescence signal were measured within 6 μm diameter ROIs encompassing the basolateral end of selected hair cells. GCaMP fluorescence intensity was calculated relative to the mean baseline intensity of each ROI prior to drug exposure.

**Flow Cytometry Sorting of Hair Cells.** Double transgenic 5 dpf larvae (-lmyosin6b:ribeyea-mcherry + neurod:GFP) were anesthetized with 0.2 mg/ml of tricaine, then transferred to preheated 37°C TrypLE (1X; ThermoFisher). Cells were dissociated at 37°C for 20 minutes with occasional trituration using a glass Pasteur pipette. The cells were spun down in a refrigerated microfuge at low speed (2000 RPM, 3 min.), then resuspended in sorting buffer (1X PBS, 2% Fetal Bovine Serum, 4 mM EDTA, 20 μM EDTA, 20 μM EDTA, 20 μM EDTA). Cells were strained through 35 μm nylon mesh (Falcon/Corning) and sorted at low pressure using a BD FACSAria IIu cell sorter; mCherry- and GFP-positive cell populations, and negative – expressing cell populations, and collected in tubes containing lysis buffer (RNeasy Micro Kit; Qiagen) for subsequent RT-PCR.

**RT-PCR.** Total RNA was extracted from isolated cells using the RNeasy Micro Kit (Qiagen). Reverse transcription (RT)-PCR was performed using RNA to cDNA EcoDry™ Premix Double Primed (Takara). Primers used for each transcript are as follows – Controls: myo6b, 5'-GCGAAGCTTGAGGTCGTCT-3' and 5'-CAGCGGAGCAAAATTGTCTT-3'; yst1a, 5'-AGTGGCATCATGGGATCATT-3' and 5'-ATCCGTGACGC-3'; 5'-actin, 5'-AGATGACAGATCATGTTGAC-3' and 5'-GTTCGATACTGCTT-3'; AMPA receptor subunits: gria1a, 5'-CAGCGGCAAACCTGCGACA-3' and 5'-CATGGCCAGCCACACC-3'; gria1b, 5'-TTGAGAATCTGAAGATTGTC-3' and 5'-CGCTCTGTCTGTGTCTCTCA-3'; gria2a, 5'-TCGAAATGCTGAGAAGACTG-3' and 5'-GCTCTCTGATGTACTGAC-3'; gria2b, 5'-GACGGACCTTTGAC-3' and 5'-ACCCCTAAATAGGGCTGTA-3'; gria3a, 5'-AGGGACCTCCTGAGAAT-3' and 5'-ACCTTCAACATATGAGGCTGTA-3'; gria3b, 5'-TGCGAGACTGTG-3' and 5'-CCACATTCTTGGCCGATAAGGA-3'; gria4a, 5'-GACTTCACACCCAGGTCTCTGTC-3' and 5'-CCACATTCTTGGCCGATA-3'; gria4b, 5'-CCATCGGAAATGCTGAACTTCT-3'; gria5a, 5'-ATGTTTCTCAGCCGACT-3'; gria5b, 5'-GGGAGACCTCAGCGACT-3'; gria6a, 5'-ATGGAGACCTCAGCGACT-3', Kainate receptor subunits: grik1a, 5'-TGCTTTGGTTGATGGCGAC-3' and 5'-AGATCCTTGACGGATTAGCC-3'; grik1b, 5'-CAGCTGTCATGGC-3'; grik2a, 5'-TGGGAAGATGGC-3'; grik2b, 5'-AGGGTGGTGGAGACTG-3'; grik3, 5'-CCAGGGCTGAGGAC-3'; grik4a, 5'-CTTACAGCCCTGCTTTC-3'; grik5, 5'-AGGGTGGTGGAGACTG-3'; grik6, 5'-CCACATTCCGCAACCGAC-3' and 5'-TGCTCAGGCACGACT-3'; grik7a, 5'-GGGAGACCTCAGCGACT-3'.
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Author Contributions
L.S. designed and performed the experiments, prepared the figures, and wrote the manuscript.

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