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Live imaging the phagocytic activity of inner ear supporting cells in response to hair cell death

EL Monzack¹, LA May¹, S Roy¹, JE Gale² and LL Cunningham^{*,1}

Hearing loss and balance disorders affect millions of people worldwide. Sensory transduction in the inner ear requires both mechanosensory hair cells (HCs) and surrounding glia-like supporting cells (SCs). HCs are susceptible to death from aging, noise overexposure, and treatment with therapeutic drugs that have ototoxic side effects; these ototoxic drugs include the aminoglycoside antibiotics and the antineoplastic drug cisplatin. Although both classes of drugs are known to kill HCs, their effects on SCs are less well understood. Recent data indicate that SCs sense and respond to HC stress, and that their responses can influence HC death, survival, and phagocytosis. These responses to HC stress and death are critical to the health of the inner ear. Here we have used live confocal imaging of the adult mouse utricle, to examine the SC responses to HC death caused by aminoglycosides or cisplatin. Our data indicate that when HCs are killed by aminoglycosides, SCs efficiently remove HC corpses from the sensory epithelium in a process that includes constricting the apical portion of the HC after loss of membrane integrity. SCs then form a phagosome, which can completely engulf the remaining HC body, a phenomenon not previously reported in mammals. In contrast, cisplatin treatment results in accumulation of dead HCs in the sensory epithelium, accompanied by an increase in SC death. The surviving SCs constrict fewer HCs and display impaired phagocytosis. These data are supported by *in vivo* experiments, in which cochlear SCs show reduced capacity for scar formation in cisplatin-treated mice compared with those treated with aminoglycosides. Together, these data point to a broader defect in the ability of the cisplatin-treated SCs, to preserve tissue health in the mature mammalian inner ear.

Cell Death and Differentiation (2015) 22, 1995–2005; doi:10.1038/cdd.2015.48; published online 1 May 2015

Hearing loss affects more than 360 million people worldwide and is often irreversible.¹ Mechanosensory hair cells (HCs), the receptor cells of hearing and balance, are not regenerated in the adult mammal and their death results in permanent hearing loss.^{2,3} HCs are surrounded by glia-like supporting cells (SCs) that are necessary for HC survival and function (reviewed in Monzack et al.).⁴ SCs perform many functions, including providing critical trophic factors, preventing excitotoxicity, and mediating regeneration in those systems (non-mammalian vertebrates) capable of replacing lost HCs.5-11 When HCs die, SCs also preserve the integrity and function of the remaining tissue by forming scars and clearing dead HCs.^{2,12-17} Maintaining a fluid barrier at the surface of the sensory epithelium after damage is necessary to preserve the electro-chemical gradient that drives HC depolarization and therefore sensory transduction after the onset of hearing (reviewed in Wangemann).¹⁸

Several major stressors cause HC death,^{19–22} including aging, noise trauma, and exposure to therapeutic drugs with ototoxic side effects. When a HC is killed by noise or aminoglycoside antibiotics, surrounding SCs form a filamentous actin (F-actin) cable that constricts the HC at its apex.^{2,12–17} This process separates the apical portion of the cell, including the stereocilia bundle, from the HC body and preserves a sealed reticular lamina.²³ In the chick utricle, following the apical constriction of dead HCs, the SCs engulf and phagocytose the remaining HC corpse.¹⁵ Additional data from the chick indicate that the ototoxic drug cisplatin impairs some SC functions, including regeneration of HCs or clearance of HC debris.²⁴ We hypothesized that SCs would have significant phagocytic activity in the mature mammalian inner ear, and that cisplatin would impair this activity. To examine these dynamic processes, we live-imaged SC phagocytic activity in the adult mouse utricle and compared the SC responses with HC stress and death caused by aminoglycosides *versus* cisplatin.

Results

Loss of tdTomato signals death. Our live-imaging experiments use whole-organ utricle cultures from adult transgenic mice that express tdTomato in all HCs. In initial experiments, dying HCs consistently lost tdTomato fluorescence and we examined whether loss of tdTomato is a reliable indicator of HC death. Live-imaging experiments were conducted with TOTO-3 iodide, a plasma membrane-impermeant dye and a dead cell indicator.^{15,25} Figure 1a shows a field of tdTomato-positive (red) HCs cultured in the aminoglycoside antibiotic neomycin as imaging begins, with the subsequent

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¹National Institute on Deafness and Other Communication Disorders, National Institutes of Health, Bethesda, MD, USA and ²UCL Ear Institute, University College, London WC1X 8EE, UK

^{*}Corresponding author: L L Cunningham, National Institute on Deafness and Other Communication Disorders, National Institutes of Health, Building 35A Room 1D971, 35A Convent Drive, Bethesda, MD 20892, USA. Tel: +1 301 443 2766; E-mail: lisa.cunningham@nih.gov

Abbreviations: ABR, auditory brainstem response; F-actin, filamentous actin; HC, hair cell; MIP, maximum intensity projection; ROI, region of interest; SC, supporting cell; SPB, Sorensen's phosphate buffer

Received 24.11.14; revised 20.2.15; accepted 04.3.15; Edited by M Piacentini; published online 01.5.15

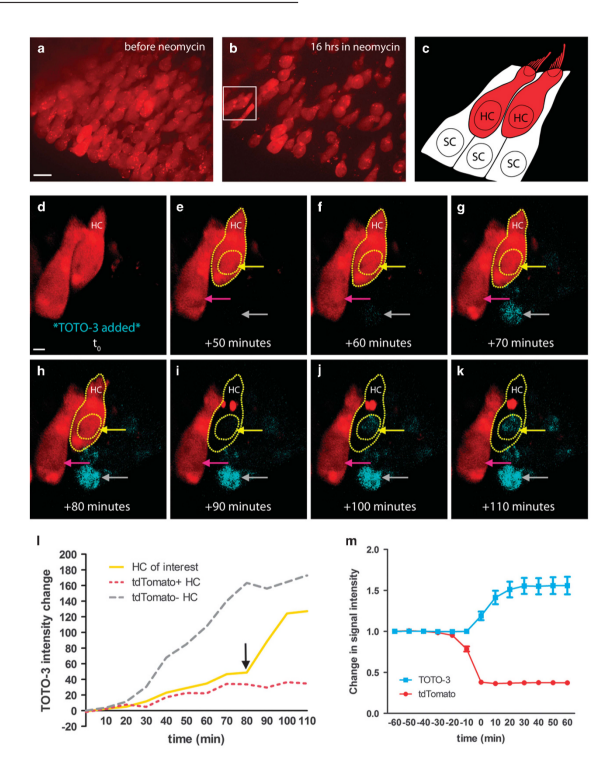


Figure 1 Loss of tdTomato signals HC death. (a) HCs (red) at the onset of live imaging before the addition of neomycin. Maximum intensity projection (MIP), scale bar = 10 μ m. (b) Culture in neomycin for 16 h results in death of many HCs. Box indicates the HCs that are shown at higher magnification in d–k. (c) Schematic of the boxed region from b. Two HCs are visible in the center of the field. SCs completely surround each HC in the utricle, but are not expressing any fluorophore and are therefore not visible in the image series. (d–k) TOTO-3 was added to cultures as a dead cell indicator, following 19 h of imaging in neomycin (t₀). Gray arrow indicates a TOTO-3-positive HC nucleus that lost membrane integrity before the addition of TOTO-3. Scale bar = 2 μ m. (h and i) The HC outlined in yellow (HC) loses tdTomato signal between +80 and +90 min. (i and j) The TOTO-3 signal in this nucleus increases over the next 20 min of imaging, becoming strongly visible. The nucleus from the tdTomato-positive HC that does not die in this image series does not show an increase in TOTO-3 (magenta arrow). (I) The mean TOTO-3 fluorescent intensity of each indicated nucleus (above) was quantified and its change over time was plotted. The black arrow marks the last tdTomato-positive time point of the HC of interest. (m) In a separate set of experiments, TOTO-3 signal intensity was quantified for 60 min before and after HC death (loss of tdTomato) in neomycin. On average, cells became tdTomato-negative and TOTO-3-positive in the same 10-min imaging interval. Movie available as Supplementary Figure 1. N = 26-28 HCs across 6 image fields from 2 utricles per time point; values represent mean ± S.E.M.

loss of HCs evident after 16 h (Figure 1b). The inset in Figure 1b highlights two HCs further analyzed following the addition of TOTO-3. Figure 1c illustrates the orientation of these two HCs and schematizes the orientation of the surrounding SCs, which were not visible during imaging. Figure 1d (movie in Supplementary Figure 1) shows a 3D rendering of the same two HCs after 19 h in neomycin, at which point TOTO-3 was added (t_0). The HC of interest (HC, vellow outline) lost tdTomato signal between +80 and +90 min (Figures 1h and i), the same period during which TOTO-3 signal intensity increased suddenly in its nucleus (black arrow on yellow line, Figure 11). TOTO-3 signal intensity further increased over the next 10 min (vellow arrow. Figure 1i). Sixty minutes after the addition of TOTO-3 (Figure 1f), an adjacent cell nucleus that was tdTomato-negative at the time of TOTO-3 addition began to show TOTO-3 uptake (gray arrow, Figure 1f), which increased over time as TOTO-3 continued to diffuse into the tissue. During this same period, the neighboring HC showed stable tdTomato signal with no visible nuclear TOTO-3 signal (pink arrow). Figure 11 shows TOTO-3 intensity over time for each of these three cells, revealing (i) TOTO-3 diffusion into the nucleus of a tdTomatonegative HC (gray line), (ii) the sudden increase in TOTO-3 signal in the nucleus of the HC that loses tdTomato (yellow line), and (iii) the low level of nuclear TOTO-3 in the tdTomatopositive HC (pink line). HC population data show that uptake of TOTO-3 occurs in the same 10-min imaging interval as the loss of tdTomato (Figure 1m). Together, these data indicate 1997

that loss of tdTomato from the HC signals loss of membrane integrity and therefore HC death.

SCs constrict and remove HCs killed by neomycin.

To examine the SC response to HC death in the mature mammalian inner ear, we used live imaging to observe the phagocytic removal of dead HCs. Utricles were infected with Ad-LifeAct-EGFP, driving an EGFP-tagged F-actin-binding peptide in a subset of SCs.^{26,27} Figure 2 shows images selected from a live-imaging experiment in neomycin (movies in Supplementary Figures 2 and 3). One tdTomato-positive HC (red, HC1) and one LifeAct-EGFP-expressing SC (green, SC1) are labeled in Figure 2a, with the orthogonal view displayed in Figure 2a'. HC1 subsequently dies (loses tdTomato signal) at +6 h (Figures 2d and d'), followed by phagocytic activity in the neighboring SC. By +6.5 h, the green SC has enriched F-actin (arrowheads, Figures 2e and e'), constricting the apical region of the dead HC. By +15 h, the SC has formed an F-actin-rich basket-like phagosome (arrowheads, Figures 2f and f') surrounding the remaining HC corpse. The F-actin-rich phagosome has also pushed the HC corpse deeper into the sensory epithelium. Magnified inset images (Figure 2a") show a HC (HC2) and two neighboring SCs (SC2 and SC3). Following the death of HC2 at +4 h (Figure 2b"), two adjacent SCs have constricted the tdTomato-negative HC corpse within 30 min. entering the space previously occupied by the HC (Figure 2c"). The position of the tdTomato-negative HC remains relatively stable for several hours following loss of membrane integrity

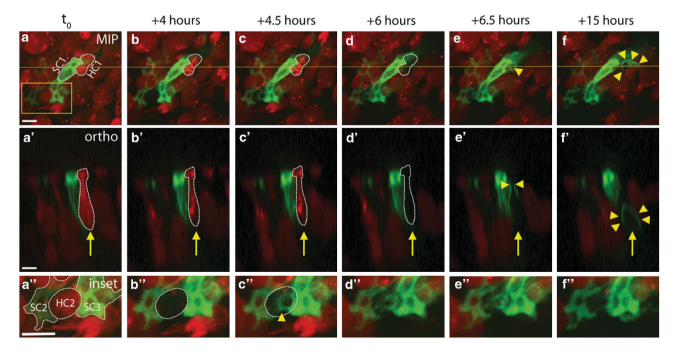


Figure 2 SCs constrict the apical portion of dead or dying HCs before removing them from the sensory epithelium. (**a**–**f**) In a series of MIPs taken from a 48-h live-imaging experiment, a single HC (red, labeled HC1 in **a**) loses tdTomato signal at +6 h (dashed outline in **d**). The neighboring green SC (labeled SC1 in **a**) is observed constricting the apical portion of HC1 within 30 min of tdTomato loss (single arrowhead in **e**). SC1 then forms a phagosome surrounding the remains of HC1, eliminating the HC corpse from the sensory epithelium (multiple arrowheads in **f**). **a**^{*}–**f**^{*} Orthogonal views of the same imaging field as **a**–**f**, taken from the plane indicated by the yellow line in **a**. Arrow follows HC1, as it loses tdTomato (dashed outline, **d**^{*}) before SC constriction (arrowheads, **e**^{*}) and phagosome formation (multiple arrowheads, **f**). **a**^{*}–**f**^{*} Magnified inset views taken from the outlined box in **a**. Two SCs (labeled SC1 and SC2 in **a**^{*}) surround a single HC (labeled HC2 in **a**^{*}), which loses tdTomato signal at +4 h (dashed outline, **b**^{*}). SC2 and SC3 together constrict HC2 within 30 min (**C**^{*}). Movies are Supplementary Figures 2 and 3. Overall gain was increased on inset images. Scale bars = 5 µm

(Supplementary Figure 4), indicating that the dead HC occupies its original space until removal by the SC phagosome.

In live-imaging experiments in which HCs were killed by neomycin, initiation of the SC response immediately followed (or occurred at a time point indistinguishable from) loss of tdTomato from the HC. To examine whether these two events were separable in time, we decreased the imaging interval from 30 to 10 min. Figure 3a shows three views from the same image field (movies are Supplementary Figure 5). Figures 3a–i are 3D surface views of a red HC (dashed outline) and its neighboring green SC. Figures 3a'–i' are differential maximum intensity projections (MIPs), in which the change in signal relative to the previous frame is shown in white. Figures 3a"–i" show a 3D orthogonal view of the same area. In this series the HC loses tdTomato (dashed outline, Figures 3b, b' and b"), and within 10 min the SC has formed an F-actin-based process in direct apposition to the apical portion of the tdTomato-negative HC (arrowhead in Figure 3c"). Thirty minutes later, the SC processes have advanced toward the center of the dead HC and formed an apical constriction (arrowheads, Figures 3f and f"). Although we cannot pinpoint the moment of bundle excision due to the loss of tdTomato in the HC, the apical constriction appears to be tightly closed in Figures 3h–3h", with the phagosome structure visible 10 min later (arrowheads in Figure 3i"). Our data indicate that SCs first move to constrict neighboring HCs either in the same frame as HC death or later (Figure 3j). On average, SCs constricted the dead HC within 1 h of tdTomato loss (Figure 3k).

To examine the contents of the SC phagosome, TOTO-3 was used in longer-term live-imaging experiments. Figure 4 shows a series of 3D reconstructions (movie is Supplementary Figure 8) in which a HC dies (dashed outline, Figure 4c) and TOTO-3 enters the nucleus (arrowheads in Figure 4c, strongly

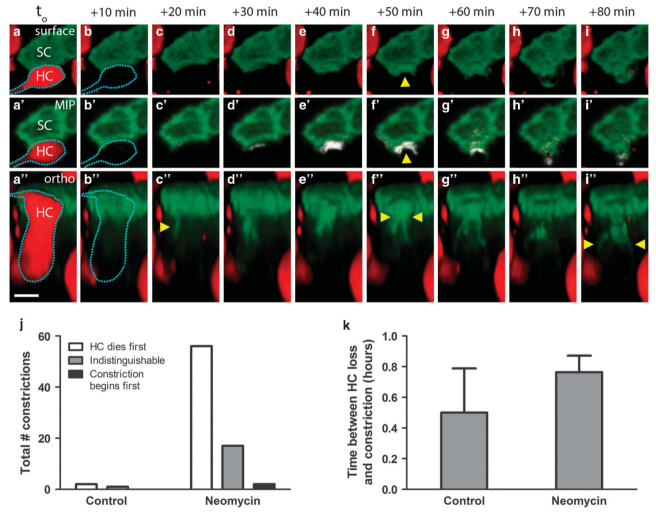


Figure 3 SC constriction begins shortly after HC death. (a–i) 3D surface views of a single SC (SC, green) and neighboring HC (HC, red). (a'–i') MIP view of the same image field, containing a third channel (white) that displays the change in EGFP signal from the previous frame. a''–i'' 3D orthogonal view of same HC and SC. The HC loses tdTomato signal (dashed outline in **b**–b'') and the SC initiates constriction toward the HC within 10 min (arrowhead, c''). Thirty minutes later, the apical constriction is visible (arrowheads, f–f''), followed by the phagosome structure (arrowheads, i''). Scale bar = 5 μ m. (j) Initiation of SC phagocytic activity occurs after or concurrent with HC death. Bars represent the sum total of all constrictions counted across four to five utricles per condition. (k) When SCs constricted dead or dying HCs, the time between the loss of HC membrane integrity and initiation of SC constriction of the apical portion of the HC averaged <1 h, regardless of the culture condition. Movies are Supplementary Figure 5. N = 3-75 SC constrictions across 4–5 utricles per condition. Bars represent mean \pm S.E.M.

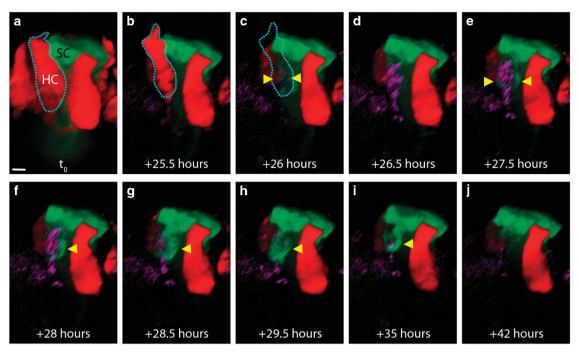


Figure 4 The SC phagosome contains the nucleus of a dead HC. (**a** and **b**) 3D orthogonal view of a red HC and an adjacent green SC. (**c**) During live imaging, the HC loses tdTomato signal in the same frame in which TOTO-3 is weakly visible (arrowheads). (**d**) Thirty minutes later, the TOTO-3 is easily visible within the HC nucleus. (**e**): Within 1.5 h of tdTomato loss, the green SC has extended processes to surround the TOTO-3-positive nucleus (arrowheads). (**f**) A phagosome begins to surround the nucleus (arrowhead). (**g** and **h**) The SC phagosome has fully surrounded the nucleus. (**i** and **j**) The HC has been degraded and the phagosome appears to have receded. Movie is Supplementary Figure 8. Scale bar = 3.5 μ m

visible in Figure 4d). Ninety minutes after HC death, the adjacent SC first extends processes toward the HC corpse (arrowheads, Figure 4e), which then further extend and fully surround the TOTO-3-positive nucleus as a phagosome (arrowhead, Figures 4f–h). The phagosome then gradually recedes (Figures 4i–j). These data confirm that the SC phagosome contains a dead HC, and that SCs are actively removing HC corpses from the sensory epithelium.

SC phagocytic activity is reduced in cisplatin-treated utricles. We also examined SC phagocytic activity when HCs were killed by the other major ototoxic drug, cisplatin. The time courses of HC death are different for neomycinversus cisplatin-treated utricles: the average time required for 50% of the HCs to die was 23 ± 6 h in neomycin and 45 ± 3 h in cisplatin (mean \pm S.D.). To compare across conditions such that we were examining utricles with similar numbers of dead HCs, we treated with each drug for 48 h and performed subsequent quantitative analyses relative to the point when $50 \pm 5\%$ of the HCs had died (lost tdTomato signal). Neomycin-treated utricles exhibited a significant increase in the number of SC constrictions relative to controls. However, when utricles were treated with cisplatin, there was no increase in constrictions (Figure 5a). In addition, when 50% of the HCs had died, those cultured in neomycin did not show any SC death, while those cultured in cisplatin exhibited nearly 50% SC death (Figure 5b).

After the stereocilia bundle is excised, SCs form actin-rich scars at the surface of the epithelium.^{2,12–17} We examined scar formation in fixed utricles after treatment with either neomycin or cisplatin. These utricles were not live-imaged but

were cultured in neomycin or cisplatin until 50% of the HCs died, given an additional 4 h for the maximal SC response, and were fixed and stained for F-actin. In both control and neomycin-treated utricles, SC scars were consistently observed where HCs were missing (Figure 5c). In contrast, cisplatin-treated utricles revealed many dead HCs that were not covered by SC scars, a defect which persisted when utricles were cultured longer in cisplatin to allow for additional HC death and more time for SCs to respond (Supplementary Figure 9). Together, these data indicate that SCs are capable of robust scar formation in response to neomycin-induced HC death, while cisplatin-treated SCs show impaired scar-forming ability and an increase in cell death.

HC corpses accumulate in cisplatin-treated utricles.

As our experiments indicate that cisplatin impairs scar formation, we hypothesized that cisplatin treatment would also impair the clearance of HC corpses by SCs. We cultured utricles in neomycin or cisplatin until 50% of the HCs died and then allowed an additional 4 h for HC clearance. We then fixed the tissue and stained for myosin VIIA, a commonly used immunochemical marker that is specifically expressed in HCs.^{28-32} We consistently observed tdTomato-negative (dead) HCs that showed immunoreactivity for myosin VIIA (Figure 6a). As our data indicate that loss of tdTomato is a reliable indicator of HC death, these tdTomato-negative/ myosin VIIA-positive HCs are corpses. Figure 6a shows control, neomycin-treated, and cisplatin-treated utricles stained for myosin VIIA. In merged images at the HC nuclear level (Figure 6a, merge), asterisks indicate all HC corpses. Merged images from the epithelial surface, the reticular

lamina of the same image stacks (Figure 6a, final column) show the positions of the neighboring SCs (F-actin, white) relative to the myosin VIIA and tdTomato channels (individual channels, Supplementary Figure 10). HC corpses increased modestly in neomycin-treated utricles over controls. However, HC corpses in cisplatin-treated utricles increased significantly relative to neomycin-treated tissue (Figures 6a and b). This difference in corpse accumulation is not attributable to a neomycin-induced reduction in myosin VIIA expression, as there was no difference in myosin VIIA signal intensity between living (tdTomato+) and dead (tdTomato-) HCs cultured in either neomycin or cisplatin (Figure 6c). Together, these data indicate that cisplatin impairs SC-mediated phagocytic removal of dead HCs.

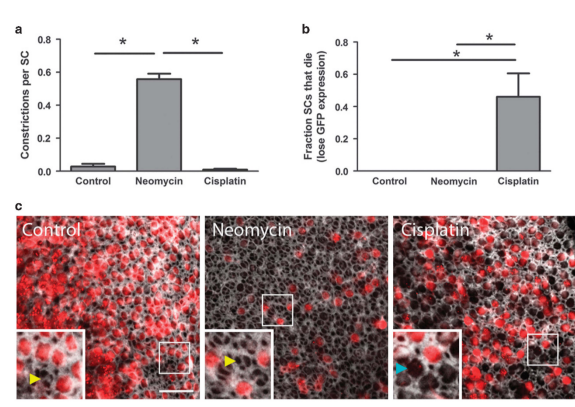
HCs shrink before death. In addition to the differences we observed between cisplatin- and neomycin-treated tissue, we also observed similarities. We quantified HC volume in the hours before death (Figure 6d). In either neomycin or cisplatin, HCs exhibited similar decreases (>150 μ m³) in cell volume before losing tdTomato. In addition, there appears to be an increased rate of volume loss in the final 3–4 h before HC death. These data indicate that the HCs undergo a cell death program that begins hours before loss of membrane

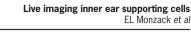
integrity, regardless of the ototoxic drug that induces HC death.

Cisplatin impairs the SC response to HC death in vivo.

To investigate whether our ex vivo data reflect the mature mouse cochlea in vivo, we examined the SC responses to HC death in cochleas from adult mice that received systemic aminoglycosides or cisplatin. Figure 7 shows 3D renderings of F-actin (labeled with phalloidin) in cochleas from mice that were treated with saline, the aminoglycoside antibiotic kanamycin, or cisplatin. Cochleas from saline-treated mice showed normal cochlear architecture with three rows of outer HCs and a single row of inner HCs (Figure 7a). Cochleas from both kanamycin- and cisplatin-treated mice showed significant loss of outer HCs. Cochleas from kanamycintreated mice showed thick, F-actin-based scars at the reticular lamina (Figure 7b). However, the scars from cisplatin-treated mice were thinner and less robust (Figure 7c). Quantitative analysis of phalloidin signal intensity indicated that scars formed in response to cisplatin treatment contained significantly less F-actin than scars formed in response to aminoglycoside treatment (Figure 7d). These data indicate that the SC responses to HC death are also impaired by cisplatin in the mouse cochlea in vivo.

Figure 5 Cisplatin impairs SC phagocytic activity. Utricles were treated with either neomycin or cisplatin until 50% of the HCs died. (a) SCs in neomycin-treated utricles exhibit robust phagocytic activity, as indicated by the number of SC constrictions. This activity is significantly reduced when HCs are killed by cisplatin. N = 3-75 SC constrictions across 4–5 utricles per condition. *P < 0.05 versus neomycin. (b) When 50% of the HCs have died, no SCs have died in control or neomycin-treated utricles; however, cisplatin-treated utricles exhibit a significant increase in SC death (determined by complete loss of EGFP signal). Bars represent mean \pm S.E.M. N = 126-276 SCs across 4–5 utricles per condition. *P < 0.05 versus control. (c) In a separate set of experiments, utricles were not live-imaged, but were cultured until 50% of the HCs died, then fixed and stained with phalloidin (white). In a single plane taken beneath the stereocilia bundles, scars (yellow arrowheads, inset) are visible in both the control and neomycin-treated utricles in spaces where tdTomato-positive HCs (red) are missing. In cisplatin-treated utricles, spaces missing tdTomato-positive HCs were often not covered by SC scars (blue arrowhead, inset). Scale bar = 20 μ m. Overall gain was increased on all inset images





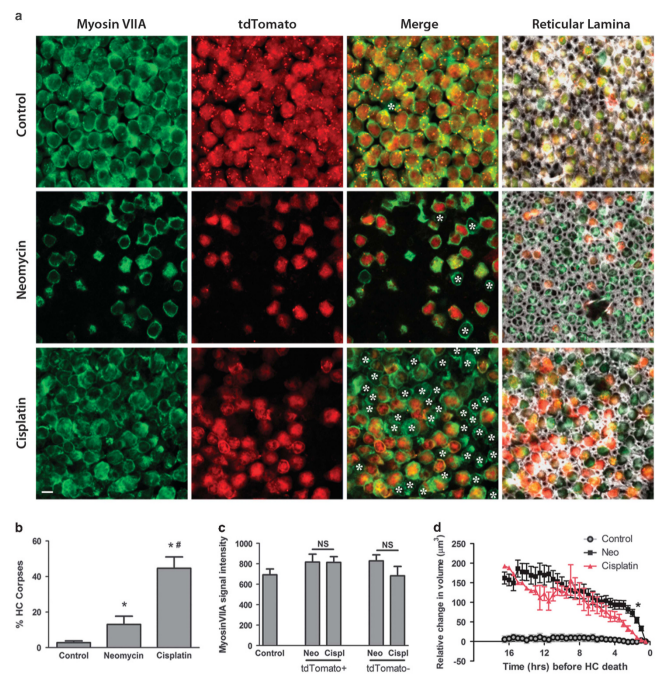


Figure 6 HC corpses accumulate in cisplatin-treated utricles. (a) Utricles were cultured in either control medium (upper row), neomycin (middle row), or cisplatin (bottom row) until 50% of the HCs died (or a matched time point in control medium). Four hours later, utricles were fixed and stained for the HC marker myosin VIIA (green), which was compared with the tdTomato signal present in living HCs (red). HC corpses (cells that are myosin VIIA-positive and tdTomato-negative) are indicated by asterisks in merged images from each condition. All images were acquired at the plane of the HC nuclei, with the exception of those in the fourth column, which were taken at the level of the reticular lamina and include the f-actin signal (white) in the merged image. Scale bar = 5 μ m. (b) The percentage of HC corpses was significantly increased in neomycin-treated utricles over controls, while the percentage of HC corpses was highest in the cisplatin-treated utricles. Bars represent mean \pm S.E.M. N = 358-2583 HCs across 3–4 utricles per condition. *P < 0.05 versus control, $^{#}P < 0.05$ versus neomycin. (c) Myosin VIIA signal intensity was not reduced in neomycin-treated tissue relative to cisplatin-treated tissue. Both tdTomato- HCs were analyzed at the nuclear plane. Bars represent mean \pm S.E.M. N = 7 HCs per condition. (d) HC volume was analyzed over time in control, neomycin, and cisplatin-treated tissue. As the HCs cultured in either neomycin or cisplatin approached HC death (indicated by loss of tdTomato signal at t = 0 h), HCs from both conditions shrank in volume, particularly in the last 3–4 h before death. HCs cultured in control medium were tracked for the same amount of time as the drug-treated utricles, but they did not die and did not shrink in volume. N = 1-31 HCs per time point, values represent mean \pm S.E.M. *P < 0.0002 via *t*-test with Bonferroni correction, neomycin versus cisplatin



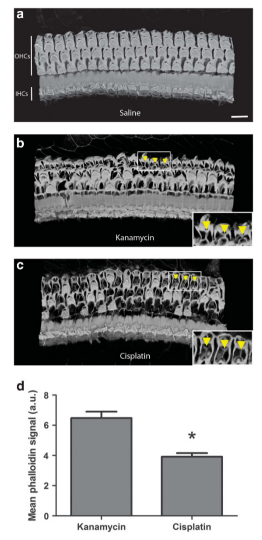


Figure 7 Cisplatin results in defects in SC phagocytic activity in mouse cochlea *in vivo.* (a) In cochleas taken from a saline-treated mouse and stained with phalloidin to label F-actin, a single row of inner HCs (IHCs) and three rows of outer HCs (OHCs) are present and have intact stereocilia bundles. Scale bar = $14 \,\mu$ m. (b) In cochleas of mice treated with systemic kanamycin, many dead OHCs have been replaced with actin-rich SC scars (a subset of which is highlighted by arrowheads). (c) In contrast, when mice were treated with cisplatin, the scars that replaced missing OHCs were thinner and composed of less actin (arrowheads) than those formed in cochleas of kanamycin-treated mice. Insets show magnified scars from OHC regions (arrowheads). (d) F-actin signal intensity was reduced in scars from cisplatin-treated mice relative to those of kanamycin-treated mice. N = 36-43 scars across 3 cochleas per condition. **P*<0.05 *versus* kanamycin-treated mice

Discussion

Our work examined the SC response to HC death caused by both major classes of ototoxic drugs. When HCs are killed by aminoglycoside antibiotics, our data are consistent with previous studies showing that SCs constrict the apical portion of HCs.^{13–15,17,33,34} Phagosome formation has previously been reported in immature avian utricles¹⁵ and our study is the first to use live imaging to examine the formation of basket-like, F-actin-rich phagosomes in the mature mammalian inner ear. We followed individual HCs over time, as they lost membrane integrity, internalized TOTO-3, and were encapsulated and

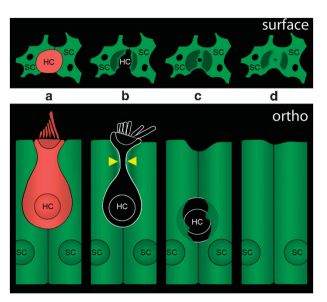


Figure 8 Model of SC phagocytic activity following HC death. (a) The architecture of a normal utricle includes SCs surrounding a healthy HC (surface view in top panel, orthogonal view in bottom panel). (b) Following treatment with aminoglycoside antibiotics, SCs constrict the HC beneath the cuticular plate (yellow arrowheads). Loss of tdTomato from the HC indicates loss of HC membrane integrity and death. (c) SC constriction results in excision of the stereocilia bundle and cuticular plate from the HC. SCs extend processes to form a phagosome around the remains of the dead HC. (d) In the final stage of the process, the HC corpse has been removed and the SCs have formed a scar at the reticular lamina

degraded by neighboring SCs. TOTO-3 labeled dead HCs following the loss of tdTomato, confirming that SC phagosomes contain nuclei of tdTomato-negative HC corpses. These data are supported by previous reports of HC nuclei or debris within SCs after damage.^{15,35–37}

Our data are consistent with a model in which glia-like SCs remove dead HCs from the mature mammalian inner ear (Figure 8). SCs completely surround healthy HCs (Figure 8a). Following exposure to an ototoxic aminoglycoside antibiotic (i.e., neomycin), HCs lose membrane integrity (lose tdTomato) and SCs constrict dead HCs beneath the cuticular plate (Figure 8b, arrowheads). This constriction excises the stereocilia bundle and cuticular plate from the HC, while SCs further constrict the apical portion of the dead HC and extend F-actin processes to form a phagosome around the HC corpse (Figure 8c). Our data suggest that the SCs ultimately remove the remains of the dead HC and form a scar at the reticular lamina (Figure 8d). Our experiments using TOTO-3 indicate that the dead HC nucleus remains fairly static, enclosed within an SC phagosome for several hours. This is a slower time course of engulfment and removal than was described in the chick15 and it is consistent with observations regarding the enhanced responsiveness of chick versus mammalian SCs during wound healing.³⁸ We also observed instances in which SCs ultimately push the HC corpse deeper into the epithelium, consistent with the idea that macrophages may also participate in the late stages of removal of dead HCs.³⁹

SCs generally advanced toward the apex of the HC concurrently with or following HC death. This is in contrast with what has been reported in the developing chick, in which SCs constricted the apical portion of the HC before loss of

membrane integrity and TOTO-3 uptake occurred concurrently with phagosome formation.¹⁵ The SC constriction in the embryonic chick was also faster than was observed here in the mammal. In the chick utricle, the constrictive event lasted \sim 180 s, as opposed to the 30-min (or longer) constrictions we often observed in the adult mouse utricle. However, as ~ 1/3 of the constrictions in our study began in the same imaging frame in which the HC lost tdTomato, it was not possible to conclusively determine which event occurred first in those instances. This close timing between the loss of tdTomato and the constriction by the SC suggests that SCs were likely responding to HC death, possibly signaled by the release of intracellular ATP.4,40,41 It is also possible that HCs prime their surrounding SCs with a stress signal before they lose membrane integrity. The idea that HCs may signal their impending death is supported by our observation that dying HCs shrink for several hours before losing membrane integrity, a phenomenon that occurred in both neomycin- and cisplatintreated utricles. These data suggest that HC shrinkage occurred independently of SC activity, consistent with recent data reported in the mammalian cochlea.35 The slow shrinkage that precedes HC death may thus represent the initiation of a cell death program that began hours before any SC activity was observed.

Our data indicate that the SC phagocytic response, which is robust in aminoglycoside-treated tissue, is nearly absent in cisplatin-treated tissue. This suggests that either (1) the stress signal emitted from dying HCs is different in response to aminoglycosides versus cisplatin treatment or (2) the SCs themselves (or at least their phagocytic responses) were impaired by cisplatin. The number of apical constrictions was reduced to nearly zero in cisplatin-treated tissue. This finding, coupled with an increase in SC death, implies that SCs are impaired by cisplatin but not by aminoglycosides. Although the cisplatin-induced death of ~ 50% of SCs would be expected to reduce SC activity, this loss does not account for the near-total loss of constrictions we observed in cisplatin-treated tissue. The surviving cisplatin-treated SCs were almost never observed constricting the apical portion of a HC, despite being surrounded by the same amount of HC death as the neomycin-treated SCs. The idea that cisplatin specifically impairs SC activity is reinforced by previous work showing that cisplatin increased apoptosis in dissociated chick SC cultures, and it impaired regeneration and HC debris removal in the chick utricle.²⁴ In addition, a recent study indicates that cisplatin (but not neomycin) damages cell-cell junctions ⁴² and another study reported deficient scar formation in the adult mouse utricle upon treatment with Latrunculin A, which prevents actin polymerization.¹⁷ The resulting pattern at the reticular lamina reported in Burns and Corwin¹⁷ is reminiscent of that reported here following cisplatin treatment (Figure 5c), consistent with cisplatin disrupting actin dynamics.43-45 Our in vivo data show that phalangeal scars formed in aminoglycoside-treated cochleas are more actin-rich than those formed in cisplatin-treated cochleas (Figure 8), further suggesting that cisplatin impairs F-actin dynamics in the inner ear.

We observed an accumulation of HC corpses within cisplatin-treated sensory epithelia. This corpse accumulation is consistent with both the lower level of SC phagocytic activity we observed and the increase in SC death. Accumulation of HC corpses has further implications for tissue health, as uncleared apoptotic cells would likely release proinflammatory intracellular contents into the surrounding tissue, causing secondary necrosis.⁴⁶ In addition, our observation of tdTomato-negative/myosin VIIA-positive HC corpses highlights a caveat of using immunochemical HC markers, as a percentage of the labeled cells are likely to be corpses.

Data from our labs and others indicate that SCs have vital roles in determining whether HCs under stress ultimately live or die. The current study indicates that these cells are critical after HCs die. Reminiscent of the role microglia have in removing neuronal debris,⁴⁷ SCs remove HC corpses and preserve the remaining health of the tissue. However, important questions remain about intercellular signals between dying HCs and their surrounding glia-like SCs, the role of resident macrophages, and exactly how long in advance of its death the HC begins to send 'distress signals'. Understanding intercellular signaling between these cell types will enhance our understanding of the roles of SCs and their responses to HC stress and death.

Materials and Methods

All reagents were obtained from Sigma Aldrich (St Louis, MO, USA), unless otherwise specified.

Utricle cultures. All mice were obtained from the Jackson Laboratory (Bar Harbor, ME, USA). The offspring resulting from a cross between B6.Cg-Tg (Atoh1-cre)1Bfri/J and B6.Cg-Gt(*ROSA*)26Sor/^{tm14}(CAG-tdTomato)Hze/J strains were used for all *in vitro* experiments. The mice were euthanized by C0₂ asphyxiation followed by decapitation; all animal procedures were approved by the NINDS/NIDCD Animal Care and Use Committee. The adult mouse utricle preparation has been previously described in detail.^{26,48} Briefly, temporal bones were removed from adult Atoh1-Cre x Rosa26-tdTomato transgenic mice of both sexes (4–12 weeks old) and placed in sterile M199 (Life Technologies Corporation, Carlsbad, CA, USA). Utricles were then dissected under sterile conditions and otoconia were removed with an eyelash tool (Ted Pella, Redding, CA, USA).

For experiments that did not require live imaging, utricles were then transferred to complete culture medium (DMEM/F12 (Life Technologies Corporation)) with 50 U/ml Penicillin G and 5% fetal bovine serum (Life Technologies Corporation) overnight at 37 °C in a humidified atmosphere with 95% air/5% CO2. The following day (~20 h post dissection), utricles were treated with either 3 mM neomycin sulfate or 30 µg/ml cisplatin (APP Pharmaceuticals, LLC, Schaumburg, IL, USA). Before addition to the culture medium, neomycin sulfate was equilibrated at 37 °C, 95% air/5% CO2 for a minimum of 2 h, and cisplatin was equilibrated for a minimum of 30 min. Drugtreated utricles were cultured for the average amount of time necessary for 50% of the HC population to die, plus an additional 4 h to allow SCs time to clear dead HCs. This timing was calculated for each drug from a minimum of four separate liveimaging experiments. tdTomato-positive HCs were counted in the first frame of each experiment. When the frame contained $50 \pm 5\%$ of the original number of tdTomatopositive HCs, that time point was recorded and used to calculate the average time necessary for 50% HC death, which was 23 ± 6 h (mean \pm S.D.) in 3 mM neomycin and $45 \pm 3 h$ in μ g/ml cisplatin.

Immunostaining and HC corpse quantification. At the end of the culture period, utricles were fixed for 30–40 min in 4% paraformaldehyde in 0.1 M Sorensen's phosphate buffer (SPB) and washed in SPB. Tissue was blocked/ permeabilized for 3 h in 0.8% normal goat serum (Vector Laboratories, Burlingame, CA, USA), 0.4% Triton X-100, and 2% bovine serum albumin. Utricles were incubated overnight at 4 °C with a mouse monoclonal antibody against myosin VIIA (138-1, diluted 1:100 in blocking buffer, Developmental Studies Hybridoma Bank, Iowa City, IA, USA). This antibody was detected using Alexa Fluor 488 goat antimouse IgG (Life Technologies Corporation). Following secondary antibody incubation, utricles were stained with Phalloidin-Atto 390 for 1 h before mounting onto glass slides with Fluoromount G (Southern Biotech, Birmingham, AL, USA).

Images of the extrastriolar region within each utricle were acquired at ×40 magnification on an LSM 780 confocal microscope (Carl Zeiss Microscopy, Oberkochen, Germany).

For quantification of HC corpses (HCs that were myosin VIIA-positive and tdTomato-negative), Zen 2012 software (Blue edition, Carl Zeiss Microscopy) was used. HCs were counted in 50 × 50 µm regions of interest (ROIs), with a minimum of six ROIs examined in the extrastriolar region of each utricle. HCs were counted in these ROIs by an investigator who was blinded with respect to the tdTomato channel. Following these counts, the tdTomato channel was visually enabled and HCs that were negative for tdTomato (equivalent to background) were counted. Control utricles for each time point were quantified to determine average HC density, which was used as a reference point to ensure that each quantified area of the drug-treated utricles contained 50 \pm 25% of the control average HC density. This controlled for amount of HC death when quantifying the percentage of corpses in each area. Time-matched controls for each drug (27 h in culture to match neomycin and 49 h in culture to match cisplatin, equivalent to the mean time to reach 50% HC death in each drug-treated condition, plus 4 h for HC clearance) did not differ significantly from each other with respect to the percent of HC corpses, and so were statistically pooled into a single control condition.

In an additional set of experiments designed to maximize the chance of seeing SCs respond to HC death caused by cisplatin, utricles were incubated for additional time in cisplatin relative to neomycin. In these experiments, utricles were cultured for 20 h in 3 mM neomycin or 52 h in 30 μ g/ml cisplatin, with time-matched controls. Following fixation, utricles were stained with anti-myosin VIIA and phalloidin. Myosin VIIA signal intensity was quantified in Volocity software (version 6.3, PerkinElmer, Waltham, MA, USA) by drawing a $12 \times 12 \times 10 \,\mu$ m (66 248 voxels) ROI at the HC nuclear level, detecting objects within 1 S.D. of the mean signal, and measuring fluorescent intensity of the objects within the ROI. Twenty-hour and 52-h controls were not significantly different from each other and so all control values were pooled.

Adenovirus infection and live imaging. Adenoviral infection of SCs in adult mouse utricles has been described.^{26,28} Following dissection and otoconia removal, utricles were transferred to a Nunc MicroWell MiniTray (Thermo Scientific, Waltham, MA, USA) and infected with a recombinant adenovirus (adenovirus type 5, DE1/E3) carrying LifeAct-EGFP under the control of the human CMV promoter. Ad-LifeAct-EGFP was generated by sequencing and subcloning an ~1.0-kb *Nhel/Not*l fragment from the LifeAct-EGFP plasmid (kind gift from Guillaume Charras, UCL) into a Dual-CCM shuttle vector, which was then subcloned into an adenoviral backbone (Vector BioLabs, Philadelphia, PA, USA).

Ad-LifeAct-EGFP $(1.05 \times 10^7 \text{ plaque-forming units})$ was added to the utricles for 2 h in serum-free medium (DMEM with 50 U/ml Penicillin G). Following adenovirus infection, utricles were transferred to 24-well plates and maintained overnight in complete culture medium. For select experiments that were not used for quantification of apical constrictions, the virus-infected utricles were maintained in culture for up to 2 days before imaging.

Utricles were imaged on a spinning disk confocal microscope (Carl Zeiss Cell Observer SD) controlled by AxioVision software (Carl Zeiss Microscopy) or Zen 2012 (Black/Blue editions, Carl Zeiss Microscopy), using a Zeiss Plan-Apochromat, \times 63, 1.4 NA, oil objective that was heated to 37 °C during acquisition. Laser lines (488, 561, and 639 nm) were used in conjunction with 525/50, 617/17, and 690/50 nm emission filter sets, respectively. Images were acquired with either a Cool Snap HQ2 CCD camera (Photometrics, Tucson, AZ, USA) or an Evolve EMCCD camera (Photometrics). Imaging was performed at 10- or 30-min intervals for a maximum duration of 70 h in phenol red-free complete culture medium (control) or phenol red-free medium treated with either 3 mM neomycin or 30 μ g/ml cisplatin. All live-imaging data were obtained from the extrastriolar region. During imaging, cultures were maintained with 94% air/6% CO₂ in a humidified atmosphere.

In each figure, image frames are denoted as occurring relative to t_0 , which was selected relative to an event of interest and was a variable amount of time (0–28 h) after imaging began. Supplementary Movies and figure legends accompany each figure that included live imaging and provide details regarding the timing of each t_0 . In calculating the differential MIP in Figure 3, the image stack was cropped to the upper 5 μ m of the reticular lamina before the differential channel was created.

Loss of HC membrane integrity. To assess the timing of the loss of HC membrane integrity relative to loss of tdTomato signal, utricles were infected with Ad-LifeAct-EGFP as above and imaged at 10-min intervals for 16 h in 3 mM neomycin. After 19 h in culture, 1 μ M TOTO-3 iodide (Life Technologies Corporation) was added (delivered in 1 ml phenol red-free complete culture medium, reducing the

concentration of neomycin to 2 mM). Following TOTO-3 addition, imaging was resumed at 10 min intervals for 2 h. Nuclear TOTO-3 signal was quantified using Volocity software by measuring fluorescent intensity within a 1.74 × 1.74 × 2.25 μm (867 voxels) ROI within a selected nucleus. A blank region outside of the sensory epithelium was used to determine the background signal, which was subtracted from all intensity values.

To track TOTO-3-positive nuclei over longer periods, 0.5 μM TOTO-3 iodide was added to the culture medium at the onset of the experiment and utricles were imaged at 30-min intervals.

HC volume. HC volume was quantified using Volocity software. Measurement settings were configured to automatically detect HCs as individual objects within the tdTomato channel; the accuracy of the automatic detection was visually verified and total cellular volume was calculated for each cell at each time point. Once determined for each cell, the automatic detection threshold settings were not changed over the duration of the time series.

Quantification of SC constrictions and death. Image stacks were visually analyzed in Zen 2012 and SC constrictions were counted. The criteria used to define an SC constriction were as follows: (1) the SC's F-actin processes advanced toward or around the apical portion of the HC; (2) when the process lasted for multiple frames, there was a clear progression in movement in which the SC's processes advanced toward the center of the HC; and (3) the HC began with and subsequently lost tdTomato signal at some point during the imaging. The total number of constrictions over time (up to the point when 50% of the HCs had died) in each image set was normalized to the number of Ad-LifeAct-EGFP-infected SCs in the frame to account for any variability in SC infection efficiency. When examining the timing of the constrictions in MIPs of all movies, the beginning of the constriction was defined as the image frame in which the LifeAct-EGFP-positive F-actin processes were first observed to be extending toward or around the HC. SC death was indicated by the complete loss of EGFP signal.

In-vivo experiments. CBA/J mice of both sexes were obtained from the Jackson Laboratory and used for all *in-vivo* experiments. Mice were 2 months old at the beginning of each injection protocol, which has been described.⁴⁹ Briefly, cisplatin (APP Pharmaceuticals, LLC) was administered via intraperitoneal injection, 4 mg/kg each day for 4 days, followed by 10 days of recovery. Three such cycles of cisplatin were administered. Saline was administered in control mice. Mice were euthanized by CO₂ asphyxiation, followed by decapitation, at the end of the third cycle.

Kanamycin sulfate was dissolved in sterile saline (Hospira, Inc., Lake Forest, IL, USA), to a final concentration of 45 mg/ml. Kanamycin (750 mg/kg) was injected subcutaneously twice daily for 14 days. Mice recovered for 21 days after the final kanamycin injection. Saline was administered subcutaneously in control mice. Preand post-injection auditory brainstem response (ABR) testing was performed to examine hearing sensitivity in both cisplatin- and kanamycin-treated groups, as described.⁴⁹ ABR testing was also performed for each control group.

Preparation of whole-mount cochleas. Cochleas were fixed overnight at 4 °C with ice-cold 4% paraformaldehyde after creating a small hole at the apex and the round window. The tissues were washed in PBS and decalcified in 0.5 M EDTA for 3–4 days. Cochleas were micro-dissected into three turns (apical, middle, and basal) and stained with Phalloidin-Atto 390. Each cochlear turn was mounted on a glass slide using Fluoromount G. F-actin staining was detected using the 405-nm laser line in conjunction with the 421–490 nm emission filter set. Images(x63) of selected regions from the transition zones (areas containing both intact and missing outer HCs) of the cochlea were taken using an LSM 780 confocal microscope (Carl Zeiss Microscopy). Imaging parameters were kept constant for all comparisons across groups.

Image stacks were visualized in Volocity software and the phalloidin signal was quantified by drawing a cylindrical ROI with a diameter of $3.95\,\mu m$ and a height of $5.1\,\mu m$ (11 696 voxels). The ROIs were each placed within regions of individual missing outer HCs, centered on the reticular lamina, and fluorescent intensity within the ROI was measured.

Statistics. All statistics were performed in Prism 5 (Graphpad Software, Inc., La Jolla, CA, USA). In comparing two conditions, data were subjected to a Student's *t*-test. Experiments that required analysis of more than two conditions were

subjected to ANOVA with Tukey's *post-hoc* test, with *P*-values < 0.05 considered significant.

Conflict of Interest

The authors declare no conflict of interest.

Acknowledgements. We thank Dr Joseph Burns and Dr Jonathan Bird for their insightful comments on this manuscript. We also thank the NIDCD/NINDS animal care staff for their support during the cisplatin injection protocol, as well as Dr Jane Johnson at the University of Texas Southwestern Medical Center for generously providing the Atoh1-nGFP mice. This work was supported by the NIDCD Division of Intramural Research (ZIA DC000079) and a Flexigrant from Action on Hearing Loss.

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Supplementary Information accompanies this paper on Cell Death and Differentiation website (http://www.nature.com/cdd)