Gap Junctional Coupling is Essential for Epithelial Repair in the Avian Cochlea

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The loss of auditory hair cells triggers repair responses within the population of nonsensory supporting cells. When hair cells are irreversibly lost from the mammalian cochlea, supporting cells expand to fill the resulting lesions in the sensory epithelium, an initial repair process that is dependent on gap junctional intercellular communication (GJIC). In the chicken cochlea (the basilar papilla or BP), dying hair cells are extruded from the epithelium and supporting cells expand to fill the lesions and then replace hair cells via mitotic and/or conversion mechanisms. Here, we investigated the involvement of GJIC in the initial epithelial repair process in the aminoglycoside-damaged BP. Gentamicin-induced hair cell loss was associated with a decrease of chicken connixin43 (cCx43) immunofluorescence, yet cCx30-labeled gap junction plaques remained. Fluorescence recovery after photobleaching experiments confirmed that the GJIC remained robust in gentamicin-damaged explants, but regionally asymmetric coupling was no longer evident. Dye injections in slice preparations from undamaged BP explants identified cell types with characteristic morphologies along the neural-abneural axis, but these were electrophysiologically indistinct. In gentamicin-damaged BP, supporting cells expanded to fill space formerly occupied by hair cells and displayed more variable electrophysiological phenotypes. When GJIC was inhibited during the aminoglycoside damage paradigm, the epithelial repair response halted. Dying hair cells were retained within the sensory epithelium and supporting cells remained unexpanded. These observations suggest that repair of the auditory epithelium shares common mechanisms across vertebrate species and emphasize the importance of functional gap junctions in maintaining a homeostatic environment permissive for subsequent hair cell regeneration.

Key words: cochlea; connexin; deafness; hair cells; hearing; repair

Introduction

Vertebrate inner ear sensory epithelia contain two principal cell types: sensory mechanoreceptors known as hair cells and nonsensory supporting cells. Hair cells are damaged most commonly by noise, ototoxic drugs, and the effects of aging (Monzack and Cunningham, 2013; Rubel et al., 2013). In the inner ear of mammals, hair cell loss may lead to hearing impairment and balance disorders, but a small degree of hair cell regeneration occurs naturally in vestibular epithelia (Forge et al., 1993; Forge et al., 1998). In nonmammalian species, inner ear epithelia produce new hair cells throughout adult life, thereby continually replacing lost cells (Stone and Cotanche, 2007; Rubel et al., 2013; Wan et al., 2013). When hair cells die, supporting cells play a key role in the maintenance of epithelial barrier integrity, ensuring that fluid compartments are separated and that ion homeostasis is preserved (Gale and Jagger, 2010). Dying hair cells are eliminated rapidly and the spaces they formerly occupied are filled. These “repair” processes must be undertaken before any regenerative replacement of hair cells and recovery of function can occur. After ototoxicity in the avian cochlear epithelium (the basilar papilla or BP), some apoptotic hair cells are ejected from the epithelium (Hirose et al., 2004; Mangiardi et al., 2004) and externalized hair cell corpses are broken down via controlled signaling pathways (Mangiardi et al., 2004). In mammalian and nonmammalian inner ear epithelia, another mechanism occurs whereby adjacent supporting cells excise the dying hair cell’s apical cuticular plate and stereociliary bundle from the soma (Li et al., 1995; Gale et al., 2002; Rubel et al., 2013) and the cell body is phagocytosed by supporting cells (Forge, 1985; Abrashkin et al., 2006; Bird et al., 2010).

A damage response common between species is an apical expansion of supporting cells (Forge, 1985; Hirose et al., 2004; Abrashkin et al., 2006; Raphael et al., 2007; Taylor et al., 2008; Taylor et al., 2012). In the organ of Corti, inner hair cell lesions may be filled by inner border cells and inner phalangeal cells and outer hair cell lesions may be filled by Deiters’ cells (Taylor et al., 2012). In addition, the Deiters’ cells migrate to fill the tunnel of Corti (Taylor et al., 2012) and the columnar epithelium is subsequently remodeled as a “flat” epithelium. In the organ of Corti of
were postfix in 1% cacodylate-buffered OsO4 and then processed with reduced ejection of dying hair cells from the damaged epithelium. With reduced ejection of dying hair cells from the damaged epithelium, GJIC remained robust. The repair of connexin expression, but GJIC remained robust. The repair of connexin expression, but GJIC remained robust. The repair of connexin expression, but GJIC remained robust.

Organotypic cultures. Cochlear ducts and utricles of chicken hatchlings (Forge et al., 2013) were opened to expose the BP and loaded for 20 min with 1–2 μm calcine-AM (Invitrogen) diluted in Medium 199 with Hanks’s salt solution. After loading, culture dishes were washed several times with HBSS. Measurements were taken within the next 2–3 h on a confocal laser-scanning microscope (TCS SP2; Leica). The tissues, attached to the membrane inserts, were maintained in Medium 199 with HBSS with 25 mM HEPES at 35 ± 2°C. A 63× water-immersion apochromatic objective lens (numerical aperture 0.9) of an upright epifluorescence microscope was used to focus just below the luminal surface of the epithelium. Optical sections were scanned in xyt mode at 1.6 μm/step. A 48 μm × 48 μm square was bleached at maximum laser power for 32 s. An area of 119 μm × 119 μm was recorded for the prebleach and postbleach scans at low laser power to minimize bleaching and fluorescence recovery was monitored over 5 min. Postbleaching fluorescence intensity (I) within regions of interest (ROIs; ~7600 pixels) was expressed in 8-bit gray-level units as a function of time using LCS 2.0 software (Leica). For each sample, the instantaneous recovery rates (½t0) from six ROIs within an optical section were expressed as mean ± SD and tested for significance using the paired Student’s t-test (GraphPad Prism 4).

Connexin-null mice, the gap junctional intercellular communication (GJIC) is impaired and repair responses display peculiarities (Forge et al., 2013). Supporting cells show uncoordinated expansion and migration does not occur, implicating GJIC in normal repair. We previously characterized GJIC in supporting cells of the undamaged chicken inner ear (Nickel et al., 2006) and identified differential connexin expression coinciding with asymmetric dye coupling in which dye flowed quicker at the epithelial abneural edge than at the neural edge. Here, we investigated GJIC between BP-supporting cells during gentamicin ototoxicity. During this critical repair period, we observed dynamic changes of connexin expression, but GJIC remained robust. The repair process was impared when GJIC was blocked pharmacologically, with reduced ejection of dying hair cells from the damaged epithelium.

**Materials and Methods**

**Animals.** Fertilized eggs of white leghorn chickens (Gallus domesticus) were obtained from a national supplier (Henry Stewart and Company) and incubated at 37.4°C in a humidified, rotating incubator (Octagon 20; Brinsea) for 21 d. All data were collected from prehatch chicks of either sex at embryonic day 21 (E21). 

**Organotypic cultures.** Cochlear ducts and utricles of chicken hatchlings (E21) were dissected in cooled Medium 199 with HBSS and 25 mM HEPES. The explants were placed on membrane inserts (Millicell-PCF; Merck Millipore) and incubated in Medium 199 with Earle’s salts, 25 mM HEPES, 26 mM sodium bicarbonate, 0.69 mM L-glutamine, and 5% fetal calf serum in a 5% CO2 atmosphere at 37°C. Hair cell loss was induced by incubation in 1 mM gentamicin for 24 h. Chemicals were from Sigma-Aldrich unless stated otherwise.

**Scanning electron microscopy.** Cultures were fixed in 2.5% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.3, with 2 mM CaCl2 for 2 h. Tissues were postfixed in 1% cacodylate-buffered OsO4 and then processed through the thio-carbohydrazide-OsO4 repeated procedure (Davies and Forge, 1987) before dehydration in an ethanol series and critical point drying. They were examined in a JEOI 6700F cold-field emission instrument and digital images were collected. All digital images were adjusted for optimal contrast and brightness using Adobe Photoshop CS4.

**Antibodies.** Protein expression of chicken Cx30 was examined using a rabbit polyclonal antibody that was generated against a synthetic amino acid peptide sequence from the cytoplasmic loop of chicken Cx30 (cCx30; aa 102-112; used at 1:100), as described in our previous study (Nickel et al., 2006), and also with a commercial rabbit polyclonal antibody raised against the C terminus of mouse Cx30 (71–2200, 1:200; Invitrogen) and a commercial mouse monoclonal antibody raised against the C terminus of mouse Cx30 (33–2500, 1:200; Invitrogen). CcCx30 was detected using mouse monoclonal CcCx4 antibody (C0993, 1:200; Sigma-Aldrich). A commercial polyclonal antibody raised against the amino acid peptide sequence for the cytoplasmic loop of rat Cx26 (71-0500, 1:200; Invitrogen) and a commercial monoclonal antibody raised against the C terminus of mouse Cx26 (33–5800, 1:200; Invitrogen) were used in preliminary experiments to detect cCxC26. A polyclonal antibody raised against the cytoplasmic loop of rat Cx26 (Gap 28H1, 1:400; a gift from Prof. Howard Evans, University of Cardiff) was also used in these experiments. A mouse monoclonal HCS-1 (Hair Cell Soma-1) antibody was used to label hair cells (mAb76, 1:100; a gift from Prof. J Corwin, University of Virginia).

**Immunofluorescence microscopy.** The fixed tissues (4% PFA for 30 min) were incubated in primary antibodies diluted in PBS containing 100 mM L-lysine and 0.2% Triton X-100 overnight at 4°C and then in FITC-conjugated goat anti-mouse IgG, Alexa Fluor 633–conjugated goat anti-rabbit IgG, and 1 μg/ml TRITC-conjugated phalloidin (all Invitrogen) in PBS/linsine/Triton X-100 for 1 h at room temperature. The samples were examined by confocal laser microscopy (LSM-510; Zeiss).

**Fluorescence recovery after photobleaching.** Fluorescence recovery after photobleaching (FRAP) experiments were performed and analyzed as described previously (Nickel et al., 2006). Briefly, after 24 h in culture, cochlear ducts were opened to expose the BP and loaded for 20 min with 1–2 μm calcine-AM (Invitrogen) diluted in Medium 199 with Hanks’s salt solution. After loading, culture dishes were washed several times with HBSS. Measurements were taken within the next 2–3 h on a confocal laser-scanning microscope (TCS SP2; Leica). The tissues, attached to the membrane inserts, were maintained in Medium 199 with HBSS with 25 mM HEPES at 35 ± 2°C. A 63× water-immersion apochromatic objective lens (numerical aperture 0.9) of an upright epifluorescence microscope was used to focus just below the luminal surface of the epithelium. Optical sections were scanned in xyt mode at 1.6 μm/step. A 48 μm × 48 μm square was bleached at maximum laser power for 32 s. An area of 119 μm × 119 μm was recorded for the prebleach and postbleach scans at low laser power to minimize bleaching and fluorescence recovery was monitored over 5 min. Postbleaching fluorescence intensity (I) within regions of interest (ROIs; ~7600 pixels) was expressed in 8-bit gray-level units as a function of time using LCS 2.0 software (Leica). For each sample, the instantaneous recovery rates (½t0) from six ROIs within an optical section were expressed as mean ± SD and tested for significance using the paired Student’s t-test (GraphPad Prism 4).
era and IR-DIC optics mounted on the microscope. Patch pipettes were filled with a KCl-based solution containing the following (in mM): 140 KCl, 10 NaCl, 2 MgCl₂, 5 HEPES, 0.5 EGTA, 3 Na₂ATP, and 5 glucose, pH adjusted to 7.3 with KOH. This solution was supplemented with 0.2% neurobiotin [molecular weight (MW) 287, charge +1; Vector Labs] and 0.2% Lucifer yellow (di-lithium salt; MW 443, charge −2) or 0.2% fluorescein dextran (MW 10 000, anionic; Invitrogen). These dyes are widely used in studies of gap junction permeability; some gap junctions in mammalian cochlear supporting cells display selectivity between these molecules of similar size but contrasting charge (Jagger and Forge, 2006; Taylor et al., 2012). Pipette solutions were filtered at 0.2 μm and centrifuged to remove small, insoluble particles. Pipettes had an access resistance of 2–3 MΩ, as measured in artificial perilymph. Dyes were injected via the patch electrode during 5 min whole-cell recordings. Lucifer yellow or fluorescein dextran fluorescence was imaged immediately after the experiment via the video camera. For confocal analysis, within 5 min of the termination of the recording, cells or slices were fixed in 4% PFA for 30 min at room temperature. To detect neurobiotin, slices were permeabilized (0.1% Triton X-100 for 40 min), blocked (0.1 M l-lysine, at 35°C for 40 min), and incubated for 2 h in Alexa Fluor 555–conjugated streptavidin (1:1000; Invitrogen). Measurements of supporting cell widths were performed using Zeiss LSM software and were taken at a depth of 10 μm from the luminal surface, a position that was approximately coincident with the position of hair cell nuclei in control slices.

Results

Whole-mount preparations of the BP were viable for several days in culture and retained the key cell types of the auditory epithelium, namely the hair cells and supporting cells (Fig. 1A). As shown previously for posthatch organotypic cultures (Oesterle et al., 1993; Frenz et al., 1998; Shang et al., 2010), this in vitro model could recapitulate processes of epithelial repair and regeneration that follow aminoglycoside ototoxicity in vivo. As observed in the adult chicken after chronic injections of aminoglycosides in vivo (Hirose et al., 2004; Mangiardi et al., 2004), incubation of BP cultures in 1 mM gentamicin activated hair cell death (apparent within 6 h; data not shown) and the subsequent ejection of their corpses from the sensory epithelium within 24 h (Fig. 1B). Subsequent incubation of the whole mounts in gentamicin-free medium led to the disappearance of the corpses from the epithelium’s luminal surface (Fig. 1C) concomitant with an expansion of the supporting cells and, within 3–5 d, there was the appearance of de novo hair cells identified by their characteristic immature hair bundles (Fig. 1D).

Connexin expression and dye coupling during ototoxic damage and repair

Having confirmed that the organotypic explant model provided a homeostatic environment compatible with normal epithelial repair and recovery, we next examined connexin expression during the period of gentamicin-induced damage and repair using qualitative immunofluorescence (Fig. 2). Similar to our previous observations (Nickel et al., 2006), there was a differential localization of chicken connexin43 (cCx43) and cCx30 immunofluorescence within the undamaged BP sensory epithelium imaged in transverse cryosections prepared from cultures incubated for 24 h in gentamicin-free medium (Fig. 2, control). During 24 h of gentamicin incubation, there was a progressive decrease of cCx43 expression and dye coupling during ototoxic damage and repair

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In our previous study using RT-PCR analysis and in situ hybridization, we identified a novel third connexin isoform expressed within the BP, namely cCx26 (Nickel et al., 2006). This was suggested as the avian ortholog of mammalian Cx26 based on their sequence homology of ~75%. In new experiments, however, antibodies raised to detect mammalian Cx26 did not label sections of control BP (data not shown). This lack of reactivity may be explained by nonhomology between the avian and mam-
malian orthologs in the specific regions these antibodies target. Attempts were made to generate a cCx26-specific antiserum, but these were without success.

To determine whether the decrease of cCx43-labeled gap junctions affected the kinetic properties of intercellular coupling between supporting cells, FRAP experiments were performed on BP explants after the 24 h exposure to gentamicin (Fig. 3A). As we described previously (Nickel et al., 2006), after calcein-AM loading of control cultures, there was a distinct pattern of brightly fluorescent hair cells surrounded by weakly fluorescent supporting cells. In gentamicin-treated cultures, there was a homogeneous weak fluorescence from supporting cells. After laser bleaching, there was a loss of fluorescence from hair cells and/or supporting cells within the exposed square. In control cultures, the calcein fluorescence intensity within supporting cells recovered to equilibrium within ~5 min, but not within hair cells. In gentamicin-treated cultures, fluorescence within supporting cells recovered to comparable levels, suggesting that gap junctional coupling was still operative after the ototoxic insult. We have previously reported regionally asymmetric dye transfer in calcein-loaded BP explants under normal conditions (Nickel et al., 2006). In those experiments, there was a faster fluorescence recovery rate measured in ROIs at the abneural edge of the BP compared with ROIs in the neural region. Here, asymmetric calcein transfer was again observed in control explants, but the abneural/nearl asymmetry was absent after hair cell loss (Fig. 3B).

cCx26 and cCx30 subunits assemble to form functional homomeric or heteromeric gap junction channels in vitro

Previous in situ hybridization experiments demonstrated coexpression of cCx26 and cCx30 within supporting cells of the BP sensory epithelium (Nickel et al., 2006), but we have been unable to confirm colocalization of protein expression for these subunits. As an alternative approach, we have used a heterologous expression system in which we could study cCx26 and/or cCx30 subunits in vitro to determine whether they are likely to oligomerize to form gap junctions in vivo. HeLa cells, which are naturally deficient in known connexins, can be transfected with the connexin cDNA of interest to examine their specifically (Marziano et al., 2003; Matos et al., 2007). Using this system, we tested the hypothesis that cCx26 and cCx30 can colocalize to single gap junction plaques, so they have the opportunity to form heteromeric channels as observed in the mammalian cochlea (Forge et al., 2003; Jagger and Forge, 2006). In HeLa cells transfected transiently with cDNA for cCx26-AcGFP, fluorescently tagged cCx26 was localized by confocal microscopy to large intercellular plaques and to smaller intracellular puncta (Fig. 4A). Comparable results were found for DsRed-tagged cCx30 (Fig. 4B). In cells coexpressing cCx26-AcGFP and cCx30-DsRed, gap junctional plaques labeled with either or both of the fluorescent tags were evident between some cells (Fig. 4C). In separate experiments, the intercellular plaques could be identified in live cells under epifluorescence illumination, thus allowing candidate cells to be selected for dye injection (Fig. 4D–F). In 3/4 recordings from cells expressing cCx26-AcGFP only, there was a large whole-cell membrane capacitance (30–48 pF) and subsequent intercellular transfer of Lucifer yellow (Fig. 4D), findings consistent with functional gap junctional coupling. Comparable observations were seen in 6/6 cells expressing cCx30-DsRed alone (Fig. 4E, 28–60 pF) and in 5/7 cells coexpressing cCx26-AcGFP and cCx30-DsRed (Fig. 4F, 25–88 pF). In recordings from cells without connexin-labeled plaques, the capacitance measurements were smaller (10–17 pF) and there was no Lucifer yellow transfer (4/4 recordings; data not shown). These experiments suggested that cCx26 and cCx30 can colocalize to single
Figure 4. cCx26 and cCx30 may assemble to form functional gap junctions. **A**, Postfixation confocal localization of AcGFP-tagged cCx26 expressed in HeLa cells. cCx26 is detected in intercellular plaques between adjacent cells (arrow) and within small intracellular puncta (arrowheads). Nuclei are stained using DAPI (blue). **B**, DsRed-tagged cCx30 also localizes to gap junctional plaques. **C**, In cells coexpressing cCx26 and cCx30, there are plaques double labeled with AcGFP and DsRed and also some singly labeled plaques. **D–F**, Epifluorescence videomicrographs tracking the transfer of Lucifer yellow between connexin-expressing HeLa cells after whole-cell recordings. In cells expressing cCx26 alone (**D**), cCx30 alone (**E**), or in cells coexpressing Cx26 and Cx30 (**F**), Lucifer yellow fluorescence is detected in the injected cell (*) and at least one adjacent cell. Scale bars, 10 μm.

Figure 5. Dye transfer in slice preparations of cultured BP explants. **A**, Photomicrograph of a BP slice preparation used in the experiment shown in **B** and **C** counterstained with fluorescent phalloidin to label F-actin (cyan). HoC, Homogeneous cells; NL, neural limbus; TM, tectorial membrane; TV, tegmentum vasculosum; SE, sensory epithelium. **Bi**, IR-DIC videomicrograph taken during a whole-cell patch recording from a supporting cell toward the abneural edge of the sensory epithelium. **Bii**, Epifluorescence videomicrograph showing resulting Lucifer yellow transfer between supporting cells. **Biii**, Confocal projection showing neurobiotin (red) and Lucifer yellow (green) distribution in the same slice after PFA fixation. The injected cell is denoted by an asterisk. Nuclei are stained using DAPI (blue). Cells containing neurobiotin have red/blue double-labeled nuclei. **C**, Single confocal image after a separate recording showing the distribution of neurobiotin in neural region supporting cells. Phalangeal processes extend from the cell body toward the luminal surface (arrows). Neurobiotin is not detected in hair cells. **Di**, IR-DIC videomicrograph taken during a whole-cell patch recording from a supporting cell toward the abneural edge of the sensory epithelium. **Dii, Diii**, Confocal projections showing the resulting distributions of neurobiotin (red) and Lucifer yellow (green). **E–G**, Examples of dye injections into supporting cells during transient bath application of gap junction blocker MFA (200 μM). Distinct delicate (**E** and thick (**F**)) supporting cell types are found toward the neural edge and thick cells are found toward the abneural edge (**G**) of the sensory epithelium. The phalangeal process of the thick cell in the neural region (**F**) measured 1.8 μm across. Scale bars, 20 μm.

Gap junction plaques, but the resulting (homomeric or heteromeric) channels share similar functional properties, such as their permeability to charged dye molecules. This is in contrast to observations for mammalian Cx26-containing and Cx30-containing channels, which demonstrate a differential permeability to anionic tracer molecules (Marziano et al., 2003). Indeed, the functional properties of cCx30 may be quite distinct from those of mammalian Cx30, which restricts the passage of large anionic dyes.

**Gap junctional coupling in slice preparations of the cultured basilar papilla**

The preservation of intercellular connectivity within slice preparations of the mammalian cochlea has enabled investigation of GJIC in sensory and nonsensory tissues (Jagger and Forge, 2006; Jagger et al., 2010; Kelly et al., 2011; Taylor et al., 2012; Forge et al., 2013). To investigate the properties of GJIC between avian supporting cells in situ, we have developed a novel slice preparation of the cultured BP, which presents the sensory and nonsensory tissues in a transverse orientation (**Fig. 5A**). After similar dye injection procedures to our previous studies of the mammalian cochlea, whole-cell patch-clamp recordings could be reliably performed from supporting cells observed under infrared videomicroscopy (**Fig. 5Bi**). Whole-cell recordings were performed over a timescale compatible with observations within our previous FRAP study in which recovery was complete with 5 min (Nickel et al., 2006). After the termination of the recording, careful retraction of the patch pipette enabled formation of an electrically tight membrane patch (resistance ~1 GΩ) and injected fluorescent dyes could be observed immediately via epifluorescence videomicroscopy (**Fig. 5Bi**). After brief formaldehyde fixation of the slices, confocal projections enabled tracing of both neurobiotin and Lucifer yellow within the supporting cell syncytium (**Fig. 5Bi**). The majority of supporting cells at the neural edge of the dye-filled syncytium appeared to span the baso-apical extent of the epithelium, each with a phalangeal process extending to the luminal surface and forming a calyceal structure around the hair cells. These cells likely correlate with the “thick” type of supporting cells identified in E19 chicks using clonal analysis techniques (Fekete et al., 1998). Importantly, fluorescence within single confocal images was restricted to the supporting cells only, with no signal emitting from adjoining hair cells (**Fig. 5C**). In the neural region of one slice,
the thickness of the phalangeal processes at a depth corresponding to the hair cell nuclei (measured in single confocal images at 10 μm from the luminal surface) was 1.74 ± 0.22 μm (n = 23) thick type supporting cells were also evident in the abneural region (Fig. 5D).

Consistent with the observations of intercellular calcein transfer, the gap junctions between supporting cells close to the neural edge (Fig. 5B, typical of 8 injections) and abneural edge (Fig. 5D, typical of 6 injections) of the epithelium were permeable to both neurobiotin tracer and Lucifer yellow. This is in contrast to observations in mammalian cochlea, in which gap junctions between Deiters’ cells allow passage of neurobiotin, but not of Lucifer yellow (Jagger and Forge, 2006; Taylor et al., 2012). There was, however, no transfer of the high-MW fluorescein dextran (MW 10 000; data not shown). Additional evidence that the observed intercellular transfer of neurobiotin and Lucifer was mediated via gap junctions was obtained from experiments using pharmacological blockers. Calcein transfer via BP gap junctions is blocked by carbenoxolone (Nickel et al., 2006). In preliminary patch-clamp experiments in the present study, we found that, although bath applied 100 μm carbenoxolone did block GJIC evident as decreases of intercellular conductance and membrane capacitance, its effects were irreversible on washout. Comparable inhibition of junctional conductance and capacitance could also be achieved by meclofenamic acid (MFA), another recognized gap junction blocker (Pan et al., 2007; Kelly et al., 2012; Toycieh et al., 2013), but this block was reversible on washout and subsequently repeatable. When recordings were performed in the presence of 200 μM MFA, there was no intercellular spread of either neurobiotin or Lucifer yellow (Fig. 5E–G), confirming the effects of MFA on GJIC. The restriction of dyes to the injected cell enabled closer inspection of the morphology of individual supporting cells. Some cells at the neural edge often spanned the whole baso-apical extent of the epithelium in a manner similar to thick supporting cells, but these cells had a cell body region close to the basilar membrane and a distinctive fine cytoplasmic process extending apically to a head structure within the hair cell region (Fig. 5E). These cells likely correlate with the “delicate” supporting cells described previously and proposed as putative precursor cells (Fekete et al., 1998). The characteristic morphology of thick supporting cells could also be delineated clearly when gap junctions were blocked (Fig. 5F,G).

Functional and morphological changes in supporting cells during ototoxic damage and repair

Dye injections in supporting cells from gentamicin-damaged BP explants yielded similar results to those from control cultures, with comparable intercellular transfer of neurobiotin and Lucifer yellow (Fig. 6A). The dye filling revealed that supporting cells had expanded after the loss of hair cells, particularly in the apical region. This was a consistent observation in various locations within the BP (Fig. 6B–D). Whereas the supporting cells in an untreated slice had thin phalangeal processes (Fig. 5C), in a corresponding region of a slice from a gentamicin-treated BP, the supporting cells were significantly wider (4.63 ± 0.84 μm, n = 16, t < 0.0001, unpaired t test). This apical expansion was demonstrated in individual cells when gap junctions were blocked transiently using 200 μM MFA (Fig. 6E–G). The restricted dye spread in the presence of MFA also delineated subcellular features such as a “foot” structure, connecting the cell body to the basilar membrane (Fig. 6E–G). After some experiments, despite the presence of MFA, the dyes were apparent in adjacent cell pairs, clearly revealed by the neurobiotin-staining of two near-adjacent nuclei (Fig. 6H,1). In these injections, there was usually one supporting cell that spanned the distance between the basilar membrane and the luminal surface of the epithelium and a second, smaller cell located at the luminal border. In the example shown in Figure 6I, the smaller cell was directly overlying the larger cell within the slice and confocal projection could not easily distinguish the pair (Fig. 6Ii,iiv). However, orthogonal projections more clearly defined the cells’ morphology and the spatial relationship between them (Fig. 6iiii,iiv).
weakly rectifying $K^+$ currents (1 mM $Na^+$, $n = 2$), chloride channels (100 $\mu M$ DIDS, $n = 2$; 100 $\mu M$ 9-AC, $n = 2$), and TRP channels (100 $\mu M$ 2-APB, $n = 2$) were all without effect. However, in 6/6 cells, the conductance decreased during whole bath replacement of $Na^+$ ions with N-methyl-D-glucamine (NMDG) partially reduced the conductance of gentamicin-treated cells.

GJIC is essential for normal epithelial repair

Because functional GJIC appears to be essential for normal repair in the mammalian cochlea (Forge et al., 2013), we investigated whether gap junctions contribute to epithelial repair in the avian cochlea (Fig. 8). In transverse cryosections prepared from undamaged BP cultured with 200 $\mu M$ MFA added to the medium for 24 h before fixation (control), HCS-1 antiserum identified hair cells within a single layer at the luminal border of the sensory epithelium (Fig. 8A). In surface preparations of control explants, phalloidin labeled F-actin within the hair cell stereociliary bundles, but also at the intercellular borders, where supporting cells contact hair cells (Fig. 8B). These results suggest that chronic block of GJIC in itself does not influence hair cell survival in vitro. In gentamicin-damaged explants, HCS-1-labeled hair cell corpses aggregated within a mass outside the sensory epithelium between the luminal surface (delineated by actin labeling) and the tectorial membrane (Fig. 8C) and the apical region of supporting cells had expanded to fill the space left by dying hair cells (Fig. 8D). However, in explants cultured for 24 h in both 1 mM gentamicin and 200 $\mu M$ MFA, the hair cells were not ejected from the epithelium (Fig. 8E,F) even though their nuclei appeared condensed (Fig. 8E, inset), consistent with the hair cells having begun the process of apoptosis. Separate dye injection experiments were performed in slice preparations from explants incubated in the presence of both gentamicin and MFA for 24 h (Fig. 9). Supporting cells in these explants did not have the expanded phenotype observed previously in explants treated with gentamicin only (Fig. 6). Instead, they more closely resembled the thick and delicate morphologies described in undamaged BP. The phalangeal processes of seven thick cells measured 1.5–2.3 $\mu m$ across, a similar range to measurements in undamaged BP. The nuclei were not located luminally. At the same level of the unexpanded phalangeal processes, pyknotic hair cell nuclei were evident, confirming that dying hair cells were not fully ejected from the epithelium. Nonsteroidal anti-inflammatory drugs such as MFA are known to also inhibit cyclooxygenase (COX) isozymes, accounting for some of their therapeutic actions (Kalgutkar et al., 2000). To discount the likelihood that MFA was exerting its effects via this mechanism in our experiments, we coincubated explants with 1 mM gentamicin and the alternative COX inhibitor diclofenac (100 $\mu M$). In these control experiments, hair cells were

During the bath application of MFA, whole-cell recordings from supporting cells allowed isolation of voltage-gated currents normally masked by the gap junctional conductance (Fig. 7). Supporting cells from gentamicin-damaged BP could be described broadly as having a low membrane conductance (typically 1–5 nS measured between $-20 \text{ mV}$ and $+20 \text{ mV}$; Fig. 7A, left) or high conductance (typically $>20 \text{ nS}$; Fig. 7A, right). Supporting cells from undamaged BP displayed the low conductance phenotype only (Fig. 7B) regardless of their thick or delicate morphological classification. Low conductance cells from both conditions had membrane currents that activated and deactivated quickly and showed little voltage dependence. These properties were reminiscent of currents in supporting cells of the pigeon semicircular canal (Masetto and Correia, 1997), but gap junctional conductances were not reported in those cells. In the higher conductance cells, the voltage-activated currents displayed a slow inactivation and activation during hyperpolarizing and depolarizing steps, respectively. Attempts to identify the membrane channels mediating the large currents were unsuccessful. Blockers of

Figure 7. Divergent electrophysiological phenotypes of supporting cells in gentamicin-damaged BP. A, Whole-cell patch recordings during transient application of 200 $\mu M$ MFA from individual supporting cells in a slice preparations from gentamicin-damaged BP explants. One neural edge supporting cell (dye injection is shown in Fig. 6D) had a membrane conductance $<2$ nS (left, slope conductance measured between $-20 \text{ mV}$ and $+20 \text{ mV}$). Under the same conditions, another neural edge supporting cell (shown in Fig. 6E) had a membrane conductance $>100$ nS (right). In both recordings, Lucifer yellow fluorescence was restricted to a single cell, confirming a lack of intercellular coupling. B, Slope conductance measurements from supporting cells in control and gentamicin-treated BPs. There was a larger spread of conductance measurements from the gentamicin-treated cells, with high conductance values in cells from all locations along the neural-abneural axis. C, Whole-bath replacement of $Na^+$ ions with N-methyl-D-glucamine (NMDG) partially reduced the conductance of gentamicin-treated cells.
This suggests that decreased prostaglandin synthesis was unlikely to be the cause of the observed MFA-associated effects on epithelial repair. Cultures coincubated with 1 mM gentamicin and 200 μM 18-β glycyrrhetinic acid, another recognized gap junction blocker (Pan et al., 2007), did not undergo hair cell ejection (data not shown), further implicating GJIC with normal repair of the BP.

Discussion
Ototoxicity-associated regulation of epithelial connexins and gap junctional coupling

Previously, our studies have resolved properties of supporting cell gap junctions in preparations of mammalian organ of Corti (Jagger and Forge, 2006) and chicken BP (Nickel et al., 2006) and identified connexin-dependent processes during epithelial repair after the loss of hair cells from the mouse cochlea (Taylor et al., 2012; Forge et al., 2013). We have now observed changes of connexin expression during ototoxic insult to the chick BP, measured concomitant alterations to the properties of intercellular dye coupling, and confirmed a role for GJIC in processes mediating the ejection of dying hair cells from the damaged epithelium. These observations expand our understanding of how gap-junction-dependent signaling contributes to normal tissue homeostasis and confirms its importance in the coordination of sensory tissue repair across different species.

By screening inner ear tissues, we previously identified chicken orthologs to the mammalian Cx30, Cx26 and Cx43, as major connexin isoforms in the avian inner ear (Nickel et al., 2006). In situ hybridization suggested that, analogous to the mammalian inner ear, cCx26 and cCx30 transcripts were expressed in sensory and ion-transporting epithelia of the cochlear duct and utricle. Furthermore, their overlapping mRNA expression pattern suggested possible colocalization within individual inner ear gap junctions similar to that found in the mammalian inner ear. We attempted to producing an antibody targeted specifically toward chicken cCx26 have been fruitless thus far, so we have not been able to demonstrate colabeling of single gap junction plaques by cCx26 and cCx30 definitively in vivo. Heterologous expression suggested that cCx26 and cCx30 are able to colocalize within single gap junction plaques, but any resulting heteromeric cCx26/Cx30 gap junction channels are unlikely to share the properties of equivalent mammalian channels, particularly because Cx30 appears to be less selective than its mammalian counterpart (Marziano et al., 2003). In the BP, we demonstrated decreased expression of Cx43 in response to gentamicin-induced hair cell loss and concluded that the properties of Cx30 alone could support the observed gap junctional communication occurring during epithelial repair. The decreased expression of Cx43 may be permissive for the increased proliferative capacity of the damaged BP (for further discussion, see Nickel et al., 2006).

Avian inner ear gap junctions are permeable to large, anionic fluorescent tracers such as calcine and Lucifer yellow. This is in contrast to the hearing mammalian cochlea, in which the presence of heteromeric Cx26/Cx30 gap junction channels impairs the transfer of large anions between the supporting cells adjacent to the sensory hair cells (Jagger and Forge, 2006; Taylor et al., 2012). In addition, in the organ of Corti, there appears to be considerable local restriction of intercellular neurobiotin spread, suggesting compartmentalization of the supporting cells adjacent to the different hair cell types. This may be the basis of proposed distinct medial and lateral K⁺ recirculation pathways (Spicer and Schulte, 1998; Jagger et al., 2010). In the avian cochlea, the pathways of neurobiotin transfer appeared to be less restricted or compartmentalized. We have observed comparable unrestricted neurobiotin coupling in the adult mouse utricle (R. Taylor and D. Jagger, unpublished data), where the supporting cells may also be relatively unspecialized.

The specific localization of Cx43 to abneural BP supporting cells seems linked to the asymmetric dye coupling observed normally between those cells, a specialization not seen in the chick utricle, where Cx43-labeled puncta are not observed (Nickel et al., 2006). The apical expansion of BP supporting cells in response to hair cell loss appeared to be associated with Cx30-dependent GJIC. An equivalent mechanism is interrupted in Cx30-null mice, in which the bodies of gap-junction-deficient Deiters’ cells fail to fill the spaces left by dying outer hair cells and also subsequently fail to migrate during formation of the “flat” epithelium that remains after all hair cells have died (Forge et al., 2013). How connexin-mediated signaling coordinates supporting cell expansion and corpse removal remains open to speculation, but it may be important to study...
mechanisms such as gap-junction-mediated Ca\(^{2+}\)-waves and the subsequent activation of Ca\(^{2+}\)-dependent processes that regulate cell volume and motility.

**Heterogeneity of supporting cells in the avian auditory sensory epithelium**

It is not clear whether spatial or temporal separation of BP regenerative responses results from heterogeneity of supporting cells or plasticity among a homogeneous population. Although the majority of molecular markers for supporting cells tested so far are expressed in identical. Previous studies have identified stem-cell-like characteristics of supporting cells may not be

Gap junction inhibition during ototoxic damage prevents supporting cell expansion. Whole-cell dye injections of neurobiotin (red) and Lucifer yellow (green) into separate supporting cells close to the neural edge of a slice preparation from a BP cultured 24 h in 1 mM gentamicin and 200 μM MFA reveal unexpanded cell bodies and phalangeal processes. Pyknotic nuclei of hair cell corpses are evident within the epithelium (arrows). One cell (SC1) retains a delicate phenotype, with a very thin cytoplasmic stalk rising from the nucleus (located near the basilar membrane) to an apical head structure at the luminal surface. The extracellular recording medium was supplemented with 200 μM MFA. The slope conductance of this cell was \(-100\; nS\). The other supporting cell (SC2) with a thick phenotype has a phalangeal process (measuring 1.8 μm across) projecting from the cell body and its nucleus is located within the midsection of the sensory epithelium. The whole-cell slope conductance of this cell was \(-1\; nS\). Scale bar, 20 μm.

Morphologically, we identified thick and delicate cells near the neural edge in undamaged BP, but these were indistinguishable electrophysiologically. In gentamicin-treated explants, we identified a subset of supporting cells that had higher membrane conductance (typically 30–100 nS) and membrane potential close to 0 mV. There was no clear correlation between the occurrence of the higher conductance and a particular morphological supporting cell type. Although this conductance was not identified, we speculate that it may indicate the activation of physiological intracellular signaling pathways involved with supporting cell expansion, so it deserves further study.

**Tissue homeostasis, regeneration, and cell turnover in the damaged avian inner ear**

An essential aspect of normal tissue homeostasis is the replacement of dead or damaged cells. Two distinct mechanisms maintain the cellular integrity of a tissue: cell turnover in response to a continuous loss of differentiated cells and regeneration after a pathological stimulus that causes the death or damage of a cell population. It is now well established that, after hair cell loss in the nonmammalian inner ear, supporting cells give rise to new hair cells by two distinctive mechanisms: direct trans-differentiation and cell division (Roberson et al., 2004; Taylor and Forge, 2005). In the chicken BP, the principal model for hair cell regeneration in warm-blooded vertebrates, supporting cells are quiescent in the undamaged sensory epithelium but, in response to hair cell loss, auditory supporting cells reenter the cell cycle, proliferate, and subsequently differentiate into new functional hair cells (Stone and Cotanche, 2007).

In contrast to the BP, the postembryonic vestibular epithelia produce hair cells continuously, which is thought to occur as a result of spontaneous apoptosis of hair cells (Warchol et al., 2001).

BP supporting cells enter the S-phase 12–24 h after the onset of a damaging stimulus and proliferation reaches its maximum within 2–3 d before returning to predamaged levels after 5–7 d (Stone et al., 1999). Supporting cells give rise to either hair cells or supporting cells through symmetric or asymmetric cell division (Stone and Rubel, 2000; Stone and Cotanche, 2007). In our dye injection experiments on gentamicin-damaged BP in which gap junctions were blocked transiently, there were incidences of dye-filled cell pairs. These pairs had nuclei located toward the luminal surface of the epithelium and, although their apical pole extended toward the luminal surface, one cell of the pair appeared to remain in contact with the basilar membrane. We suggest that these observations reflect the late stages of asymmetric cell division activated by the ototoxic insult (Stone and Cotanche, 2007; Brignull et al., 2009; Wan et al., 2013), with the smaller of the pair possibly representing a newly regenerated hair cell precursor. An alternative possibility is that these cell pairs are exclusively coupled via MFA-resistant gap junctions. Future studies using our slice preparation approach to further characterize the mechanisms of hair cell regeneration could incorporate an analysis of markers such as Atoh1, which is expressed in dividing supporting cells or newly postmitotic cells (Cafaro et al., 2007). We also observed cells apparently mediating endocytic removal of hair cell corpses (Bird et al., 2010), suggesting that homeostatic repair mechanisms endure vibratome sectioning of explants. The slice preparation should prove to be a powerful new tool in the study of damage and repair mechanisms in the avian inner ear.
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