Lis1 mediates planar polarity of auditory hair cells through regulation of microtubule organization

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SUMMARY
The V-shaped hair bundles atop auditory hair cells and their uniform orientation are manifestations of epithelial planar cell polarity (PCP) required for proper perception of sound. PCP is regulated at the tissue level by a conserved core Wnt/PCP pathway. However, the hair cell-intrinsic polarity machinery is poorly understood. Recent findings implicate hair cell microtubules in planar polarization of hair cells. To elucidate the microtubule-mediated polarity pathway, we analyzed Lis1 function in the auditory sensory epithelium in the mouse. We show that conditional deletion of Lis1 in developing hair cells causes defects in cytoplasmic dynein and microtubule organization, resulting in planar polarity defects without overt effects on the core PCP pathway. Lis1 ablation during embryonic development results in defects in hair bundle morphology and orientation, cellular organization and junctional nectin localization. We present evidence that Lis1 regulates localized Rac-PAK signaling in embryonic hair cells, probably through microtubule-associated Tiam1, a guanine nucleotide exchange factor for Rac. Lis1 ablation in postnatal hair cells significantly disrupts centrosome anchoring and the normal V-shape of hair bundles, accompanied by defects in the pericentriolar matrix and microtubule organization. Lis1 is also required for proper positioning of the Golgi complex and mitochondria as well as for hair cell survival. Together, our results demonstrate that Lis1 mediates the planar polarity of hair cells through regulation of microtubule organization downstream of the tissue polarity pathway.

KEY WORDS: Lis1, Pafah1b1, PCP, Rac GTPase, Dynein, Microtubule, Stereociliary bundle, Nectin, Deafness

INTRODUCTION
In addition to apical-basal polarity, planar cell polarity (PCP), or polarity within an epithelial cell sheet, is crucial for epithelial morphogenesis and function. Sensory hair cells are specialized neuroepithelial cells that convert mechanical stimuli into electrical nerve impulses. Mechanotransduction is accomplished by the V-shaped stereociliary bundle (or hair bundle) located on the apex of each hair cell. Each hair bundle consists of rows of actin-based stereocilia arranged in a staircase pattern with a single microtubule-based kinocilium next to the tallest row of stereocilia. This polarized structure renders the hair bundle directionally sensitive to deflections (Kazmierczak and Müller, 2012). As a result, hair cells must be uniformly oriented across the plane of the auditory sensory epithelium, the organ of Corti, such that the vertex of each bundle points toward the lateral edge of the cochlear duct. Proper hair bundle polarity and orientation are essential for normal hearing.

Hair cell planar polarity is established during embryogenesis in vertebrates. One of the earliest manifestations of planar polarization is the migration of the axonemal kinocilium from the center of the hair cell apex toward the cell periphery (Denman-Johnson and Forge, 1999; Tilney et al., 1992). The kinocilium is a specialized primary cilium that extends from the basal body located just below the apical surface. Following its migration toward the lateral edge of the cochlear duct, microvilli adjacent to the kinocilium thicken and elongate, eventually forming a V-shaped bundle of stereocilia, with the kinocilium at its vertex. Nascent hair bundles then refine and align their orientation during late embryonic and early postnatal development (Dabdoub et al., 2003). Thus, positioning of the kinocilium/basal body near the lateral pole is tightly coupled with hair bundle polarity and orientation, and together they constitute morphological features of hair cell planar polarity.

Hair cell planar polarity is coordinated at the tissue level by an evolutionarily conserved core Wnt/PCP pathway (Goodrich and Strutt, 2011). Downstream of the core PCP genes, the small GTPase Rac and its effector p21-activated kinases (PAKs) have been shown to mediate basal body positioning and hair bundle orientation (Grimsley-Myers et al., 2009; Sipe and Lu, 2011). Wnt/PCP signaling spatially orients localized Rac-PAK signaling on the hair cell cortex (Grimsley-Myers et al., 2009); however, the underlying mechanisms remain incompletely understood.

Accumulating evidence suggests that tissue-level PCP signaling impinges on a hair cell-intrinsic pathway that controls the planar polarization of individual hair cells. The kinocilium and its connection to the adjacent stereocilium, via the kinociliary links, are required for the normal V-shape and orientation of the nascent hair bundle (Jones et al., 2008; Sipe and Lu, 2011; Webb et al., 2011). Moreover, a non-ciliary function of Kif3a, a component of the kinesin II plus-end-directed microtubule motor complex, coordinates basal body positioning and hair bundle orientation through spatial regulation of Rac-PAK signaling, thus implicating microtubule-based intracellular transport in hair cell planar polarization (Sipe and Lu, 2011). In addition to templating the kinocilium, the basal body (or the mother centriole), along with the daughter centriole and the associated pericentriolar matrix, organize cytoplasmic microtubules in hair cells (Steyger et al., 1989).

To further elucidate the microtubule-mediated hair cell polarity pathway, we investigated the function of a well-established microtubule regulator, Lis1 (Pafah1b1 – Mouse Genome Database).
**MATERIALS AND METHODS**

**Mice**

Animal care and use were in compliance with NIH guidelines and the Animal Care and Use Committee at the University of Virginia. Mice were obtained from the Jackson Laboratory or the referenced sources and maintained on a mixed genetic background. The morning of a plug was designated as embryonic day (E) 0.5 and the day of birth as postnatal day (P) 0. For embryonic experiments, Atoh1Cre; Lis1flox/– mice were mated with Lis1+/+; Cre mice to generate Lis1flox/+; Cre mice. For postnatal experiments, Atoh1Cre; Lis1flox/– mice were crossed with Lis1flox/+; Cre mice to generate Atoh1Cre; Lis1flox/+, which were served as controls. The GFP-centrin2 transgenic line (Higginbotham et al., 2004) was used to mark the centrioles. The following genotyping primers were used: 5'-AGAACCTGTAAGATGGCTG-3' and 5'-GGCTATAC-3'.

**Immunohistochemistry and image acquisition**

Otic capsules were dissected from mice of the indicated ages and fixed in 4% paraformaldehyde for 1 h at room temperature or overnight at 4°C. For dishevelled 2, dynein intermediate chain, Pcm1, Rac1-GTP, pan-Rac1, and mitochondrial immunostaining, otic capsules were fixed in 10% paraformaldehyde and then cryosectioned at 14 µm. Frozen in OCT (Tissue-Tek) and then cryosectioned at 14 µm.

**Quantification of hair cell phenotypes**

Quantification of hair bundle orientation was carried out as described previously (Sipe and Lu, 2011). Data in the text are presented as mean ± s.e.m. To quantify the distance of the centrosome from the cell membrane, the distance along a line drawn from the center of the centrosome to the closest edge of the cell was measured using ImageJ. For all experiments, measurements were obtained from hair cells in the basal region of the cochlea (15% cochlear length; at least three cochleae per genotype).

**RESULTS**

**Lis1 is localized to the pericentriolar region in developing hair cells**

To investigate a potential role for Lis1 in auditory hair cell development, we first determined the subcellular localization of Lis1 in the developing organ of Corti. At E17.5, Lis1 was detected on the stereocilia and the hair cell microtubule network (Fig. 1A-C). At P1, Lis1 was prominently localized to the pericentriolar region in hair cells, forming a cloud surrounding the centrosomes, and was also detected at low levels on the centrosomes of supporting cells (Fig. 1D-F). These localization patterns are consistent with those reported in other cell types and suggest a function for Lis1 in the regulation of the hair cell microtubule network (Sasaki et al., 2000; Tanaka et al., 2004).

**Lis1 deletion during embryonic development causes defects in hair bundle morphology and orientation**

To investigate the function of Lis1 in developing hair cells, we generated conditional mutants using a floxed allele of Lis1 (Hirotune et al., 1998) and an Atoh1Cre driver line that expresses Cre in developing hair cells, and a subset of supporting cells starting at ~E14.5 (Yang et al., 2010). We also used a null allele (Lis1-/-) derived from the Lis1flox allele by germline Cre expression. To perturb Lis1 function in embryonic hair cells, we generated Atoh1Cre; Lis1flox/-.
At E17.5, a subset of Lis1cKO-early hair cells had bundles with an abnormal, flattened morphology (Fig. 2B). Moreover, Lis1cKO-early hair cells displayed hair bundle misorientation (Fig. 2B,F; average bundle deviation of 20.1±1.5°) compared with littermate controls (Fig. 2A,E; average bundle deviation of 8.6±0.7°). We also examined the position of the kinocilium and found that it had migrated to the hair cell periphery (Fig. 2B). However, kinocilia were often mispositioned with respect to both the hair bundle and the medial-lateral axis of the cochlea (Fig. 2B,D). These defects in kinocilium/basal body positioning correlated with hair bundle misorientation. Furthermore, in contrast to the regular aster-shaped array in control hair cells (Fig. 2C), cytoplasmic microtubules appeared disorganized in Lis1cKO-early hair cells (Fig. 2D). These results demonstrate that Lis1 regulates the microtubule organization and planar polarization of embryonic hair cells.

Hair bundle orientation is coordinated by the core Wnt/PCP pathway (Goodrich and Strutt, 2011). Therefore, we sought to determine whether Wnt/PCP signaling is compromised in Lis1cKO-early cochlea by examining the asymmetric membrane localization of the core PCP protein dishevelled 2 (Dvl2). In E17.5 wild-type cochlea, Dvl2 was localized to the lateral side of hair cell membranes (Fig. 2G,J). This localization was essentially unchanged in Lis1cKO-early hair cells (Fig. 2H,I,K,L). Of note, the hair cell rows in Lis1cKO-early cochlea were slightly jumbled, and abnormal apical contacts between two supporting cells with Dvl2 staining were observed (Fig. 2L). These results demonstrate that Lis1 regulates the microtubule organization and planar polarization of embryonic hair cells.

Defective cellular organization and nectin localization in Lis1cKO-early organ of Corti

To further assess cellular organization in Lis1cKO-early cochlea, we stained E18.5 tissue for myosin VI and β1/β2-tubulin to label hair cells and supporting cells, respectively. Instead of the normal ‘checkerboard’ pattern of hair cells interdigitated with supporting cells (Fig. 3A), pairs of Lis1cKO-early hair cells often appeared to be in direct contact, without an intervening supporting cell (Fig. 3B). Moreover, the apical surfaces of both hair cells and supporting cells in Lis1cKO-early cochlea were misshapen. Many hair cells adopted an oblong or irregular shape and the apically spaced ‘hourglass’ shape of supporting cell apical domains was frequently jumbled and distorted (Fig. 3B). We next examined transverse cochlear sections. In control sections, hair cell nuclei were invariably located a uniform distance from the luminal surface of the epithelium (Fig. 3C). By contrast, the Lis1cKO-early sensory epithelium was disorganized, with hair cell nuclei frequently found at varying distances from the luminal surface (Fig. 3D).

The checkerboard pattern of the organ of Corti is regulated by the nectin (or poliovirus receptor-related) family of cell adhesion molecules (Togashi et al., 2011). We examined the localization of
Lis1 regulates Rac-PAK signaling in embryonic hair cells

Given the role of Lis1 as a regulator of cytoplasmic dynein in other systems, we investigated dynein localization using an antibody against the dynein intermediate chain (Dillman et al., 1994). Immunostaining of E17.5 control hair cells revealed a cloud of dynein in the apical cytoplasm, with a higher concentration in the pericentriolar region (supplementary material Fig. S2A). In Lis1cKO-early hair cells, dynein was still found in the pericentriolar region, but it was more diffuse (supplementary material Fig. S2A), indicating a role of Lis1 in regulating dynein localization in hair cells.

To further determine the mechanisms underlying the planar polarity and cellular organization defects, we next examined Rac-PAK signaling in Lis1cKO-early cochleae. Our previous work suggests a model in which microtubule-mediated transport regulates the localized activation of Rac-PAK signaling on the hair cell cortex (Sipe and Lu, 2011). This model predicts that regulators of localized activation of Rac-PAK signaling on the hair cell cortex in the control, whereas total Rac1 was uniformly distributed along the circumference of the hair cell (Fig. 4B,D,F,H). Rac1-GTP was also detected in the pericentriolar region and on the stereocilia (supplementary material Fig. S2B). In Lis1cKO-early cochleae, the cortical domain of Rac1-GTP was significantly reduced and/or misoriented relative to the medial-lateral axis of the cochlea (Fig. 4C,E,G,I). Localization of Rac1-GTP in the pericentriolar region was also disorganized and diffuse, although it was still detected on the hair bundle (supplementary material Fig. S2B). These results indicate that Lis1 regulates localized Rac1 activity at the hair cell cortex and other subcellular locations.

Rac GTPases are activated by guanine nucleotide exchange factors (GEFs) and inactivated by GTPase-activating proteins (GAPs). To investigate the molecular mechanisms of Lis1- and microtubule-mediated Rac activation, we searched the literature for Rac GEFs with reported localization to microtubules in other cell types. T-cell lymphoma invasion and metastasis 1 (Tiam1) emerged as the leading candidate. Tiam1 associates with microtubules in neurons and regulates neuronal polarity (Kunda et al., 2001). Furthermore, the Tiam1 homolog Tiam2 (or STEF) has been shown to mediate microtubule-dependent Rac activation during cell migration (Rooney et al., 2010). Importantly, cochlear explants treated with NSC23766, a competitive inhibitor of Rac interactions with certain GEFs, including Tiam1 (Akbar et al., 2006), showed planar polarity defects (Sipe and Lu, 2011). At E17.5, Tiam1 is highly enriched on microtubules and the kinocilium in control hair cells (Fig. 4J,L,N; supplementary material Fig. S2C-E). In Lis1cKO-early hair cells, although still detected on microtubules, Tiam1 staining was greatly disorganized or diminished in cells with disorganized microtubules (Fig. 4K,M,O). These data suggest a potential role for microtubule-associated Tiam1 in stimulating cortical Rac-PAK activity.

Lis1 is required to maintain the V-shape of hair bundles during postnatal development

Having demonstrated a role for Lis1 in planar polarization in embryonic hair cells, we next sought to determine the function of Lis1 in postnatal hair cells. We generated Atoh1Cre; Lis1flox/flox mice (hereafter referred to Lis1cKO-late). In contrast to Lis1cKO-early...
mutants, two Cre-mediated recombination events are required to generate a Lis1-deficient hair cell. Since Lis1 protein levels correlate tightly with gene dosage (Gambello et al., 2003), we reasoned that the combined effects of slower DNA excision and longer protein perdurance in Lis1cKO-late would enable slower depletion of Lis1 compared with Lis1cKO-early mutants.

At P0, the Lis1cKO-late organ of Corti exhibited normal cellular organization (Fig. 5B). Hair bundle morphology and orientation were also normal, suggesting that Lis1 function in the Lis1cKO-late cochlea was sufficient to sustain the embryonic phase of hair cell morphogenesis. However, starting at ~P2 and following a base-to-apex gradient along the cochlea, some Lis1cKO-late outer hair cells displayed hair bundle morphology defects ranging from a flattened bundle (Fig. 5D,F,G) to splitting of the bundle into two separate groups of stereocilia (Fig. 5D,H-J). Rarely, hair bundles were severely dysmorphic or completely fragmented (Fig. 5K). Overall, 13% of Lis1cKO-late hair bundles in outer hair cell rows exhibited a flattened morphology, 8% were split, and 1% were dysmorphic (Fig. 5E). By contrast, Lis1cKO-early vestibular hair cells in the utricular macula at P4 had no overt defects in hair bundle morphology or orientation (supplementary material Fig. S3). These results indicate that, in addition to its earlier function, Lis1 is required for maintaining the normal V-shape of the auditory hair bundle during postnatal development.

Lis1 mediates the positioning of the hair cell centrosome near the lateral cortex

To understand the basis for the hair bundle defects in Lis1cKO-late hair cells, we examined basal body positioning at P2. In control hair cells, the centrosome (the basal body and daughter centriole) was invariably found near the lateral pole and aligned along the medial-lateral axis of the cochlear duct (Fig. 5L). By contrast, the centrosome in many Lis1cKO-late hair cells was located away from the lateral pole and toward the center of the cell, particularly in those cells with a flattened or split hair bundle (Fig. 5M). Indeed, the distance between the centrosome and the lateral hair cell membrane (DC_M) closely correlated with hair bundle morphology defects (Fig. 5N). On average, DC_M in Lis1cKO-late hair cells with a normal bundle morphology (0.90±0.05 µm; n=42) was similar to that of controls (0.87±0.02 µm; n=60). However, DC_M was dramatically increased in Lis1cKO-late hair cells with a flattened hair bundle (1.55±0.06 µm; n=45) and increased still further in cells with a split hair bundle (2.04±0.07 µm; n=34) (Fig. 5N). Thus, using basal body position and the V-shaped hair bundle as morphological readouts for planar polarity, we conclude that Lis1 is required for the maintenance of planar polarity in postnatal hair cells.

To investigate whether Wnt/PCP signaling plays a role in postnatal hair cells, we examined Dvl2 localization at P2. As in embryonic hair cells, Dvl2 was asymmetrically localized on the lateral side and pericentriolar region (supplementary material Fig. S4D-I). Furthermore, Tiam1 localization to microtubules but not the cortical domain of pPAK and Pak1 in control (ctrl) and Lis1cKO-late (cKO) cochleae. Lysates of two cochleae from the same embryo were pooled and loaded in each lane. Gapdh served as loading control. Error bars indicate s.e.m. (B-I) Localization of Rac1-GTP (green) and total Rac1 (red) in E17.5 control (B,D,F,H) and Lis1cKO-early (C,E,G,I) cochleae. Boxed control and Lis1cKO-early cells are shown at higher magnification in the insets. Arrowheads mark the pillar cell row and brackets indicate outer hair cell rows. Scale bars: 6 µm.

Lis1 regulates hair cell planar polarity

Fig. 4. Lis1 mediates cortical Rac-PAK signaling, probably through microtubule-associated Tiam1. (A) Western blot analysis of the levels of pPAK and Pak1 in control (ctrl) and Lis1cKO-early (cKO) cochleae. Lysates of two cochleae from the same embryo were pooled and loaded in each lane. Gapdh served as loading control. Error bars indicate s.e.m. (B-I) Localization of Rac1-GTP (green) and total Rac1 (red) in E17.5 control (B,D,F,H) and Lis1cKO-early (C,E,G,I) cochleae. Boxed control and Lis1cKO-early cells are shown at higher magnification in the insets. Arrowheads mark the pillar cell row and brackets indicate outer hair cell rows. Scale bars: 6 µm.
Lis1 is required for proper dynein localization and centrosome organization in hair cells

To understand the basis for the centrosome positioning defects in Lis1cKO-late hair cells, we next examined the localization of cytoplasmic dynein. Immunostaining of dynein intermediate chain in control hair cells at P2 revealed a three-dimensional lattice of dynein in the pericentriolar region, extending from the centrosome basally for ~2 µm to the level of Dvl2 membrane localization (Fig. 6C). At the plane of the centrosome, dynein formed an organized ring surrounding the centrosome and also localized to one of the centrioles (Fig. 6G,J). Dynein staining was also detected on the cell cortex, partially overlapping with Dvl2 (Fig. 6E, inset). In Lis1cKO-late hair cells, instead of an organized three-dimensional lattice, dynein formed an irregular, diffuse cloud. At the plane of the centrosome, dynein collapsed inward toward the centrosome and was no longer detectable on centrioles (Fig. 6H,J). At the level of Dvl2 crescents, dynein was still detected extending basally from the pericentriolar cloud (Fig. 6D) and on the cell cortex, partially overlapping with Dvl2 (Fig. 6F, inset). These results indicate that Lis1 is required for normal dynein localization around the centrosome in hair cells.

Next we asked whether Lis1-dynein function is important for hair cell pericentriolar matrix organization, as in other cell types (Guo et al., 2006; Kubo et al., 1999; Zimmerman and Doxsey, 2000). We examined the localization of the pericentriolar material 1 (Pcm1) protein, a pericentriolar matrix component that is important for the
recruitment of other centrosomal proteins (Balczon et al., 1994; Dammermann and Merdes, 2002). In control hair cells at P3, Pcm1 was localized in a tight ring in the pericentriolar region, which formed interior to the dynein lattice ~0.2 µm below the daughter centriole (Fig. 7A,C, inset). By contrast, Pcm1 in Lis1cKO-late hair cells failed to organize into a ring-like structure and instead collapsed inward toward the centrioles, forming a diffuse cloud interior to the dynein clusters (Fig. 7B,D, inset). Thus, centrosome mispositioning in Lis1cKO-late hair cells is associated with dynein and pericentriolar matrix organization defects.

**Microtubule organization defects in Lis1cKO-late hair cells**

Lis1-dynein function in pericentriolar matrix organization regulates the interphase microtubule array in other systems (Quintyne et al., 1999; Smith et al., 2000). We therefore examined microtubule organization in Lis1cKO-late hair cells at P2, when hair bundle and dynein defects are beginning to manifest. In control hair cells, an aster-shaped radial microtubule array emanated from the pericentriolar matrix toward the cortex in all directions but was more prominent on the lateral side of the cell (Fig. 7E,G,I). Individual microtubules appeared to extend laterally and basally to contact the cortex and then curl around the circumference of the cell (Fig. 7I). In Lis1cKO-late hair cells, the organization of the cytoplasmic microtubule array was severely disrupted; the ring-like organization of the minus ends of microtubules collapsed inward and became more focused around the centrosome (Fig. 7F,H). This was particularly evident in Lis1cKO-late cells with flattened or split bundles (Fig. 7E,H,J, plane z). Instead of fanning out basally to form an aster-shaped array, microtubules in many Lis1cKO-late cells lacked apparent organization and directionality (Fig. 7F,H,J,K). In some cells, microtubules organized into thick bundles that wrapped around the circumference of the cell (Fig. 7J, plane z’). Overall, Lis1cKO-late hair cells appeared to have a reduced concentration of microtubules on the lateral side of the hair cell and more microtubules on the medial side of the cell (Fig. 7K, plane z’). Together, these results suggest that dynein and pericentriolar matrix defects in Lis1cKO-late hair cells result in disorganization of the microtubule network.

**Lis1 deficiency leads to organelle distribution defects and subsequent outer hair cell death**

In cultured cells, Lis1 and dynein regulate organelle position (Harada et al., 1998; Lam et al., 2010). We therefore determined whether the distributions of the Golgi and mitochondria were perturbed in Lis1cKO-late hair cells. At P2, the Golgi complex in control hair cells consisted of complex tubule structures (Fig. 8A) that were confined to a region just basal to the cuticular plate of the hair cell (Fig. 8B). Similarly, mitochondria were distributed throughout the cell body but were enriched in the apical cytoplasm (Fig. 8E,F). By contrast, in Lis1cKO-late hair cells, the Golgi complex appeared fragmented with vesicles spread throughout the cytoplasm (Fig. 8C,D). Mitochondria were also highly dispersed, and the apical population of mitochondria was greatly diminished (Fig. 8G,H). Taken together, these Golgi and mitochondria defects support a universal function of Lis1-dynein in organelle positioning.

To determine the fate of Lis1cKO-late hair cells, we examined Lis1cKO-late cochlea a few days after the onset of hair bundle and organelle defects. Beginning at ~P5 and proceeding in a wave from the basal region of the cochlea toward the apex, outer hair cells began to undergo apoptotic cell death, as indicated by cleaved caspase 3 immunostaining, and were extruded from the sensory epithelium (Fig. 8J). By P7, the basal region of the Lis1cKO-late cochlea was devoid of outer hair cells (Fig. 8M,N). These results reveal a requirement for Lis1 for auditory hair cell survival.

**DISCUSSION**

In this study, we have undertaken a comprehensive analysis of Lis1 and dynein function in developing auditory hair cells. Our results reveal a crucial function of Lis1 in dynein localization and microtubule organization and provide novel insights into the Lis1- and microtubule-mediated processes essential for hair cell planar polarity during both embryonic and postnatal development. In addition, we demonstrate that Lis1 is also required for proper organelle positioning and hair cell survival.

Together with recent advances, our results provide strong support for a two-tier hierarchy of hair cell planar polarity regulation. The core PCP pathway generates extrinsic or tissue polarity cues that
are interpreted within hair cells by a cell-intrinsic effector machinery. These cell-intrinsic processes are capable of operating independently of inputs from the tissue polarity pathway to drive planar polarization of individual cells, as core PCP genes are not required for establishing planar polarity features including polarized basal body position, V-shaped hair bundle and the asymmetric cortical domain of Rac-PAK activity.

We show that the asymmetric localization of Dvl2 is maintained when Lis1 is deleted predominantly in hair cells, suggesting that tissue polarity cues set up by the core PCP pathway remain intact. We propose that Lis1 is a component of the cell-intrinsic effector machinery in embryonic hair cells that controls localized cortical Rac-PAK activity through microtubule-mediated transport. At the onset of planar polarization, the arrival of the basal body at the lateral pole of the hair cell tightly correlates with asymmetric cortical PAK activity (Grimsley-Myers et al., 2009). Importantly, hair cell planar polarization is disrupted in Kif3a-deficient hair cells, where active PAK on the cell cortex is mislocalized and diffuse, indicating that plus-end-directed transport is important for constraining the cortical domain of Rac-PAK activity (Sipe and Lu, 2011). Here we show a correlation between reduced cortical Rac-PAK signaling and microtubule organization defects in Lis1l−/− hair cells, suggesting that Lis1-dependent microtubule organization is crucial for Rac-PAK activation on the cell cortex. Lis1 also appears to regulate Rac activity in the pericentriolar region, which is likely to control hair bundle cohesion and the position of the kinocilium within the hair bundle. Furthermore, we show that the Rac GEF Tiam1 is associated with microtubules in a manner that is sensitive to microtubule organization and developmentally regulated, making Tiam1 a strong candidate activator of cortical Rac-PAK signaling during the planar polarization of hair cells.

Taken together, these results suggest a model whereby microtubule-associated Tiam1 translocates to the cell cortex through Lis1-mediated microtubule-cell cortex interactions to stimulate cortical Rac-PAK activation (Fig. 9A). Migration of the basal body to the hair cell periphery is the symmetry-breaking event that sets the cell-intrinsic effector machinery in motion. Following migration of the basal body to the hair cell periphery, centriolar microtubules interact with the nearby cell cortex, allowing translocation of Tiam1 from microtubules to the cell cortex, which enables it to activate Rac-PAK signaling (Michiels et al., 1997). This cortical Rac-PAK activity strengthens local interactions between microtubules and the cell cortex through as yet unidentified cortical and cytoskeletal proteins, such as Igap1 (Kholmanskikh et al., 2006), and/or indirectly through dynein-mediated cortical capture of microtubules (Laan et al., 2012; Markus et al., 2009; Tsai et al., 2005; Yamada et al., 2008). Of note, similar mechanisms involving feedback loops between Rho family GTPases and polarized trafficking via the cytoskeleton have been proposed for self-polarization of unicellular organisms and other polarized cell types in the absence of extrinsic cues (Chang and Martin, 2009; Slaughter et al., 2009; Tahirovic and Bradke, 2009).

Acting upstream of the cell-intrinsic effector pathway, tissue-level PCP signaling may spatially coordinate cortical Rac-PAK activity through two potential mechanisms, which are not mutually exclusive. First, mechanical tension between hair cells and supporting cells may be involved in tissue-level PCP signaling. Genetic evidence suggests that the core PCP pathway acts in conjunction with a Ptk7-mediated pathway to modulate apical junctional actomyosin contractility (Lee et al., 2012; Lu et al., 2004). Anisotropic mechanical forces exerted on hair cells might bias the positioning of the basal body toward the lateral pole.
orient centriolar microtubules and align planar polarity in embryonic hair cells. In addition to actomyosin-mediated forces, the core PCP pathway may also regulate dynein-mediated microtubule capture at the hair cell cortex. Interestingly, dishevelled has been shown to regulate dynein-mediated mitotic spindle orientation through interaction with nuclear mitotic apparatus protein (NuMA) in other systems (Ségalen et al., 2010). NuMA, together with LGN (leucine-glycine-asparagine repeat-enriched protein) and Ga, forms an evolutionarily conserved cortical protein complex that recruits cytoplasmic dynein to the cell cortex to generate pulling forces that position spindle microtubules (Johnston et al., 2009; Nguyen-Ngoc et al., 2007). Recently, G protein-signaling modulator 2 (GPSM2), the human homolog of LGN, has been identified as the causative gene for the nonsyndromic deafness DFNB82 (Walsh et al., 2010; Yariz et al., 2012). Thus, we speculate that a dishevelled-dynein pathway analogous to that which mediates mitotic spindle orientation might regulate cortical microtubule capture, thereby orienting centriolar microtubules in both embryonic and postnatal hair cells (Fig. 9).

In postnatal Lis1<sup>+/KO-late</sup> hair cells, planar polarity is established normally but is subsequently lost, as indicated by the observed basal body anchoring and bundle morphology defects. This is, to our knowledge, the first direct evidence that hair cell planar polarity must be actively maintained during early postnatal development. In contrast to embryonic hair cells, cortical Rac activity is significantly downregulated in P3 hair cells, suggesting that asymmetric cortical Rac-PAK activity is crucial for the initial planar polarization process but that alternative mechanisms are employed to maintain planar polarity. Our data suggest that dishevelled might act upstream of Lis1-dynein to maintain the centrosome position in postnatal hair cells.

Intriguingly, microtubule arrays were often more tightly focused around the centrosome in Lis1<sup>+/-KO-late</sup> hair cells, consistent with loss of cortical anchoring of microtubule plus ends. In other systems, cortical dynein plays an essential role in the regulation of microtubule growth and exerts pulling forces to position microtubule-associated structures (Etienne-Manneville and Hall, 2001; Gundersen and Bulinski, 1988; Laan et al., 2012; Palazzo et al., 2001; Vallee and Stehman, 2005). The buckling and bending of long microtubules around the cell cortex observed in a subset of Lis1<sup>+/KO-late</sup> hair cells is similar to the in vitro behavior of microtubules in the absence of end-on capture by cortical dynein (Laan et al., 2012). Based on these observations, we propose that Lis1-dynein-mediated cortical capture of microtubules, together with a microtubule-organizing function at the centrosome, controls anchoring of the basal body at the lateral pole of hair cells. Proper basal body position, in turn, maintains the V-shape of the hair bundle during early postnatal development (Fig. 9B).

Although defects in dynein and microtubule organization were widespread in Lis1<sup>+/KO-late</sup> hair cells, only a subset develops flat or split hair bundles, suggesting that hair cells employ redundant mechanisms for maintaining proper hair bundle morphology. During early postnatal development, the apical region of the hair cell undergoes a shape change driven by actomyosin forces (Etournay et al., 2010). Also during this time period, densely bundled rootlet structures form at the base of the stereocilia to anchor them into the cuticular plate, an actin meshwork that provides rigid support (DeRosier and Tilney, 1989; Kitajiri et al., 2010). We suggest that the rootlets of the stereocilia and the cuticular plate serve as additional physical constraints to maintain the position of the basal body and the V-shape of the hair bundle in conjunction with Lis1-dynein. Lis1 is also required for positioning of the Golgi and mitochondria, as well as for hair cell survival. Importantly, cell death is not limited to cells with abnormal hair bundles, suggesting that it is not merely a consequence of hair bundle defects. Further study is needed to understand these additional functions of Lis1-dynein.

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References


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DEVELOPMENT

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