The planar polarity and staircase-like pattern of the hair bundle are essential to the mechanoelectrical transduction function of inner ear sensory cells. Mutations in genes encoding myosin VIIa, harmonin, cadherin 23, protocadherin 15 or sans cause Usher syndrome type I (USH1, characterized by congenital deafness, vestibular dysfunction and retinitis pigmentosa leading to blindness) in humans and hair bundle disorganization in mice. Whether the USH1 proteins are involved in common hair bundle morphogenetic processes is unknown. Here, we show that mouse models for the five USH1 genetic forms share hair bundle morphological defects. Hair bundle fragmentation and misorientation (25-52° mean kinociliary deviation, depending on the mutant) were detected as early as embryonic day 17. Abnormal differential elongation of stereocilia rows occurred in the first postnatal days. In the emerging hair bundles, myosin VIIa, the actin-binding submembrane protein harmonin-b, and the interstereocilia-kinocilium lateral link components cadherin 23 and protocadherin 15, all concentrated at stereocilia tips, in accordance with their known in vitro interactions. Soon after birth, harmonin-b switched from the tip of the stereocilia to the mid-stereocilium for the latter.

INTRODUCTION

Usher syndrome (USH) is the most frequent cause of hereditary deaf-blindness in humans. Of the three clinical forms (USH1-3), USH1 is the most severe. It is characterized by severe to profound congenital hearing impairment, constant vestibular dysfunction and pre-pubertal onset retinitis pigmentosa. Five USH1 genes have been identified. They encode the unconventional myosin VIIa (USH1B), the PDZ-domain-containing scaffold protein harmonin (USH1C), cadherin 23 (USH1D) and protocadherin 15 (USH1F), which are two cadherins with long extracellular regions, and the protein sans (USH1G), which contains ankyrin repeats and a sterile alpha motif (Fig. 1A) (Ahmed et al., 2006). The role played by these early connectors in stereocilia growth, orientation and differential growth

KEY WORDS: Usher syndrome, Hair bundle links, Planar polarity, Stereocilia growth, Harmonin (USH1C), Cadherin 23 (USH1D), Protocadherin 15 (USH1F), Sans (USH1G), Myosin VII (USH1B)
the differentiation of the hair bundle is, however, poorly understood. A scaffolding role has been ascribed to harmonin, as its PDZ domains are expected to allow the formation of large molecular complexes. Myosin VIIa is an actin filament plus-end-directed myosin, which is therefore expected to move from the base to the apex of stereocilia. Moreover, genetic evidence supports its involvement in the targeting of various hair bundle proteins, including harmonin, during postnatal stages (Boeda et al., 2002; Michalski et al., 2007; Senften et al., 2006).

Several in vitro interactions between USH1 proteins have been reported (Fig. 1C). Harmonin can bind to the four other USH1 proteins (Adato et al., 2005; Boeda et al., 2002; Reiners et al., 2005; Siemens et al., 2002; Weil et al., 2003), and myosin VIIa can interact with sans and protocadherin 15 (Adato et al., 2005; Senften et al., 2006). By contrast, only sparse evidence has been collected to support direct interactions between these proteins within the hair bundle. In addition, the precise developmental processes in which these proteins are involved are still unknown. We addressed these issues by searching for common hair bundle anomalies in mouse models for each of the five USH1 genetic forms, and by investigating the possible co-distributions and interdependent localizations of the Ush1 proteins.

MATERIALS AND METHODS

Animals and antibodies

Cdh23+/2, Pcdh15+/+; and Ush1gjgj mice were obtained from Jackson Laboratories (Bar Arbor, ME). Myo7a+/+/H9252 and Myo7a+/+ mice were kindly provided by Dr K. P. Steel (Sanger Institute, Cambridge, UK), and we produced Ush1c knockout mice (referred to as Ush1c−/−) (see Fig. S1 in the supplementary material). Embryonic day 0 (E0) was determined by vaginal plug detection, and the day of birth was considered P0. In all procedures, mice were treated in accordance with both INSERM and Pasteur Institute welfare guidelines.

Rabbit polyclonal antibodies used to detect myosin VIIa (here named Myo7a-F1; 1:1000), hair cell b (harmonin-H1b; 1:100), cadherin 23a (Cdh23-N1; 1:100) and protocadherin 15a/b (here named Pcdh15-cter; 1:500) have been described (Boeda et al., 2002; el-Amraoui et al., 1996; Reiners et al., 2005). The rabbit anti-Vangl2 antibody (1:200) was kindly provided by Drs Yao and Noda (JFCR Cancer Institute, Tokyo, Japan). Additional antibodies are commercially available: goat anti-Scrib1 (1:100; Santa Cruz Biotechnology, Santa Cruz, CA), rabbit anti-frizzled 3 (1:200; Sigma, Evry, France), mouse anti-β-catenin (1:200; Transduction Laboratories, BD Biosciences, Le Pont de Claix, France), mouse anti-acetylated-tubulin (1:200; Sigma), Alexa Fluor 488 or 546 F(ab′)2 fragment of goat anti-rabbit IgG, Alexa Fluor 488 F(ab′)2 fragment of goat anti-mouse IgG, Alexa Fluor 488 donkey anti-goat IgG (1:500; Molecular Probes-Invitrogen, Cergy Pontoise, France) and TRITC-conjugated phallolidin (1:1000; Sigma).

Whole-mount immunofluorescence

Mouse inner ears were dissected from temporal bones at different developmental stages. The cochlear shell was pierced at its apex, and the round and oval windows opened. Generally, inner ears were then immersed for fixation in 4% paraformaldehyde in PBS for 1 hour. After several washes in PBS, the cochlear shell was finely dissected and the organ of Corti processed as described (Michel et al., 2005). For Vangl2, the cochlear shell was dissected prior to fixation and the organ of Corti was only quickly fixed for 5 minutes in cold (−20°C) methanol. In this staining procedure, an anti-β-catenin antibody was used to label the apical region of hair cells because methanol has deleterious effects on the actin cytoskeleton. Fluorescence images were obtained with a confocal microscope (Zeiss LSM 510 META) equipped with a Plan Apochromat 63 X/1.4 oil immersion objective. z-stack images were deconvoluted and reconstructed with Huygens (Scientific Volume Imaging, Hilversum, the Netherlands) and Image J (Rasband WS, US NIH, Bethesda, MD; http://rsb.info.nih.gov/ij/) software, respectively.

Scanning electron microscopy

Inner ears of Ush1 mutant mice were processed as for immunofluorescence, except that they were fixed by immersion in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.3) at room temperature for 2 hours. After several washes with buffer alone, they were finely dissected to provide direct access to the cochlear and vestibular sensory areas. The samples were then dehydrated by successive washes in ethanol (50, 70, 80, 90 and 100%), critical-point dried, mounted on a stub, sputter-coated with gold-palladium and examined under a Jeol JSM6700F scanning electron microscope.

Angular deviations were measured between the expected and effective positions of the kinocilia at the apical surfaces of hair cells from the cochlear apical turn, using the graphic tools of the Jeol JSM6700F software. Briefly, for each hair cell, the kinociliary deviation was determined by the angle formed by two crossing lines. The first line was drawn mediolaterally along the symmetry/PCP axis of the cell, thereby running through the expected position of the kinocilium. The second line was traced between the center of the hair cell surface and the effective position of the kinocilium. Data were analyzed using Excel (Microsoft).

RESULTS

Among the various mouse mutants identified for each Ush1 gene, at least one carries a mutation with a predicted effect similar to that expected in the corresponding human USH1 form, except for USH1C. Indeed, the two Ush1c mouse mutants, Ush1cΔVangl2 and Ush1cΔVangl2ΔVangl2, carry mutations that affect only some harmonin isoforms (Johnson et al., 2003), whereas virtually all USH1C mutations reported so far in patients are expected to be ‘functional null alleles’ (Bitner-Glindzicz et al., 2000; Verpy et al., 2000; Zwaenepeel et al., 2001). Therefore, we engineered a harmonin-null (Ush1cΔVangl2) mouse lacking Ush1c exon 1, which contains the translation initiation site common to all reported harmonin isoforms (see Fig. S1 in the supplementary material).

Ush1 mutant mice exhibit fragmented and misoriented hair bundles from E17.5 onwards

In the mouse cochlea, postmitotic cells of the primordial sensory epithelium differentiate in concurrent medial-to-lateral (neural-to-abneural) and basal-to-apical gradients along the cochlear duct between E14.5 and 18.5. Sensory cells concomitantly become organized into one row of inner hair cells (IHCs, the genuine sensory cells) and three rows of outer hair cells (OHCs, the cochlear amplifiers) (Fig. 1B), by cell intercalations. This convergent extension process results in the shortening in width and extension in length of the sensory epithelium. Actin-rich protrusions, which will ultimately form the stereocilia, soon emerge at the hair cell apical surface (Chen and Segil, 1999; Lim and Anniko, 1985; Sher, 1971). In the mouse, temporal information regarding the differentiation of these protrusions into fully mature stereocilia is limited. In the rat and hamster, the protrusions rapidly assemble into a bundle and grow uniformly until late embryonic stages. Then, the growing stereocilia elongate differentially according to their location within the bundle, so that the “staircase-like” pattern of the stereocilia rows is settled a few days after birth. While supernumerary stereocilia are reabsorbed, the remaining stereocilia continue to grow simultaneously in length and width, until the hair bundle reaches its mature shape (Kaltenbach et al., 1994; Zine and Romand, 1996). Therefore, initial, intermediate and final periods of hair bundle development can be distinguished, in which the growth of the stereocilia row is uniform, differential and simultaneous, respectively. From our observations (see below), we established that the first two periods extend from E15.5 to P0, and from P1 to around P5, whereas the third lasts until P15, in the mouse. To determine and characterize hair bundle morphogenetic defects in
mice defective in myosin VIIa (Myo7a<sup>4626SB/4626SB</sup>), harmonin (Ush1c<sup>–/–</sup>), cadherin 23 (Cdh23<sup>v2J/v2J</sup>), protocadherin 15 (Pcdh15<sup>av3J/av3J</sup>) and sans (Ush1g<sup>js/js</sup>), we undertook a systematic and comparative analysis of the different Ush1 mutant hair bundles during these three developmental periods.

By confocal microscopy, we observed that every mutant displayed disorganized, fragmented IHC and OHC hair bundles as early as E17.5 and E18.5, respectively (Fig. 2A). Scanning electron microscopy (SEM) on P0 organs of Corti showed that in Ush1 mutants, stereocilia generally assembled into two to three clumps at the cell apical surface, instead of forming an integral, single ‘V’-shaped bundle as in wild-type hair cells (Fig. 2B). Remarkably, although lateral links that connect stereocilia to each other along their shaft were observed in stereociliary clumps of all Ush1 mutant mice, they seemed sparse compared with those of wild-type hair bundles and were also more frequently disrupted (Fig. 2C).

Examination by confocal microscopy of E18.5 cochleas stained with actin and microtubule markers revealed that a large proportion of the hair cells displayed a mispositioned kinocilium in Ush1 mouse mutants (Fig. 3A and data not shown). Moreover, observation of Ush1 mutant hair bundles by SEM at P0 showed that the stereociliary clumps were also often misoriented (Figs 2, 3).

According to the current view, the kinocilium leads the differentiation of stereocilia, which is likely to be disrupted in Ush1 mutants. This disruption could result in the mispositioning of the kinocilium and, consequently, the misorientation of stereocilia, leading to the observed phenotype. The precise mechanism by which the kinocilium is mispositioned remains to be elucidated, but it is clear that the lateral links between stereocilia are crucial for their correct positioning and orientation.

Fig. 1. The USH1 proteins and their interactions in vitro. (A) Predicted structures of the different USH1 protein isoforms. Myosin VIIa consists of a motor head, a neck region with five isoleucine-glutamine (IQ) motifs, and a large tail comprising an α-helix domain and two repeats, each composed of a myosin tail homology 4 (MYTH4) domain and a 4.1 ezrin radixin moesin (FERM) domain, separated by a Src homology 3 (SH3) domain (Chen et al., 1996; Weil et al., 1995). There are three classes of harmonin isoforms (a, b and c) (Verpy et al., 2000). Harmonin-a and harmonin-c have three and two PDZ domains, respectively, and harmonin-a also has a coiled-coil (CC) domain. Harmonin-b has the same domains as harmonin-a, plus a second CC domain and a proline, serine and threonine (PST)-rich sequence. Cadherin 23 and protocadherin 15 isoforms are also grouped into three classes (Ahmed et al., 2006; Lagziel et al., 2005). Cadherin 23a, cadherin 23b, protocadherin 15a and protocadherin 15b are transmembrane isoforms, with 27, 7, 11 and 1 extracellular cadherin (EC) repeat, respectively. Cadherin 23c isoforms are cytoplasmic, whereas protocadherin 15si isoforms are secreted. Multiple splice variants have been identified for myosin VIIa, protocadherin 15a, and each isoform class of harmonin and cadherin 23 (alternative exons are indicated) (Ahmed et al., 2006; Ahmed et al., 2003; Chen et al., 1996; Lagziel et al., 2005; Michel et al., 2005; Reiners et al., 2003; Verpy et al., 2000; Weil et al., 1995). Finally, sans has three ankyrin (ANK)-like repeats and a sterile alpha motif (SAM) domain (Kikkawa et al., 2003; Weil et al., 2003). Sans does not have any known splice variants. The respective locations of the mutations of the five Ush1 mouse models used in this study are indicated by arrows for point mutations, and by inhibition signs for the deletion (del) of the transcription start site. The resulting stop codons are indicated by asterisks. The immunogenic regions for the antibodies used in this study are indicated by brown boxes. (B) Apical views of the auditory epithelium of a P5 wild-type mouse by scanning electron microscopy (SEM). Sensory inner hair cells (IHCs) and outer hair cells (OHCs) are organized into a single medial-side row and three lateral-side rows, respectively (left panel). A hair bundle that consists of stereocilia and a single transient kinocilium is present on top of every hair cell (right). Scale bar: 1 μm. (C) Schematic representation of known in vitro interactions between the USH1 proteins. The domains involved in each interaction are drawn in close apposition. Harmonin can bind to any of the other USH1 proteins. Cadherin 23a and protocadherin 15a/b cytoplasmic regions interact with harmonin PDZ1 and/or PDZ2 domains (Adato et al., 2005; Boeda et al., 2002; Reiners et al., 2005; Siemens et al., 2002). The presence of a consensus PDZ-binding motif at the C-terminus of cadherin 23a and protocadherin 15a/b isoforms is indicated by a star. Through its cytoplasmic region, protocadherin 15a/b can also bind to the myosin VIIa SH3 domain (Senften et al., 2006). The harmonin PDZ1 domain can interact with the second MYTH4-FERM repeat of the myosin VIIa tail and the SAM domain of sans (Boeda et al., 2002; Weil et al., 2003). Finally, the central region of sans can bind to the first MYTH4-FERM repeat of myosin VIIa (Adato et al., 2005). Harmonin and sans can also form homodimers (not shown) (Adato et al., 2005).
establishment of the stereociliary bundle polarity (Jones and Chen, 2007). Therefore, we quantified the hair bundle polarity defect of the different Ush1 mutants by measuring, on each hair cell surface, the kinocilium deviation (in degrees) from its expected position along the PCP axis (arbitrarily fixed as 0°) (Fig. 3B; see Materials and methods for details). Consistent with the previous study by Dabdoub et al. (Dabdoub et al., 2003), kinocilia had mean absolute deviations ranging from 6.5° in IHCs to 12.5° in OHCs of the third row in P0 wild-type mice. They were equally distributed on both sides of the PCP axis, and more than 80% of them were present within ±15° of the PCP axis (Fig. 3C and data not shown). By contrast, most hair cells of the Ush1 mutants displayed mispositioned kinocilia. Both the IHCs and the three rows of OHCs showed larger mean kinociliary deviations than in wild-type mice (Fig. 3C). Only 37% of kinocilia at best (Ush1gjs/js), and 14% at worst (Cdh23v2J/v2J), were present within ±15° of the PCP axis in the Ush1 mutant hair cells (Fig. 3C). The mean absolute kinociliary deviations of IHCs and OHCs were 25° (Ush1gjs/js), 26° (Myo7a4626SB/4626SB), 26° (Ush1c−/−), 38° (Pcdh15av3J/av3J) and 52° (Cdh23v2J/v2J) (Fig. 3C). Nevertheless, virtually all kinocilia, apart from a small proportion in Cdh23v2J/v2J and Pcdh15av3J/av3J hair cells, were located within the lateral half and near the edge of the apical cell surface. Notably, in Cdh23v2J/v2J and Pcdh15av3J/av3J mice, the kinocilia were also often dissociated from the stereociliary clumps (Fig. 3C).

Recent studies have shown that the polarization of the kinocilium is secondary to the asymmetric distribution of core PCP proteins at cell-cell junctions of the hair cells (Deans et al., 2007). We therefore analyzed the distribution of two of these proteins, vang-like 2 (Vangl2) and frizzled 3, which are required for the establishment of inner ear PCP (Montcouquiol et al., 2003; Wang et al., 2006). In E18.5 Ush1c−/−, Cdh23v2J/v2J and Pcdh15av3J/av3J mice, Vangl2 and frizzled 3 were asymmetrically distributed on the apical, medial side of hair cells, as in wild-type littermates (see Fig. S2 in the supplementary material). In addition, scribble 1 (Scrib1; scribbled), which has also been found to be involved in inner ear PCP (Montcouquiol et al., 2003), was normally distributed along the hair cell basolateral membrane in all Ush1 mutant mice (see Fig. S2 in the supplementary material). Of note, PCP proteins are also required for cochlear convergent extension (Jones and Chen, 2007). In agreement with the normal distributions of Vangl2, frizzled 3 and Scrib1, organs of Corti in all Ush1 mouse mutants had normal length and width at P0, and they did not show extra hair cell rows (data not shown).

**Ush1 proteins localize at stereocilia tips during initial hair bundle development**

The early core phenotype that we identified in Ush1 mutant mice as early as E17.5 suggests that the Ush1 proteins cooperate during the initial period of hair bundle development. To determine whether Ush1 proteins colocalize during this period, we studied their distribution by immunofluorescence analysis of whole-mount cochleas. For each Ush1 protein, we focused on specific isoform classes that have been shown to be either prominent in, or largely restricted to, the inner ear and retina (Ahmed et al., 2006; Ahmed et al., 2003; Hasson et al., 1995; Lagziel et al., 2005; Michel et al., 2005).
2005; Reiners et al., 2003; Rzadzinska et al., 2005; Verpy et al., 2000). Thus, submembrane actin-binding class b harmonin (harmonin-b), transmembrane class a cadherin 23 (cadherin 23a), transmembrane class a and class b protocadherin 15 (protocadherin 15a/b), and myosin VIIa isoforms were specifically stained by the harmonin-H1b, Cdh23-N1, Pcdh15-cter and Myo7a-F1 antibodies, respectively (Fig. 4). The specificity of these antibodies was established by the loss of hair bundle immunoreactivity in each mouse mutant deficient for the corresponding protein (see Fig. S3 in the supplementary material). In the absence of a similar specificity indication for our anti-sans antibody, we did not study the distribution of this protein.

Myosin VIIa, harmonin-b, cadherin 23a and protocadherin 15a/b were first detected in all the emerging protrusions of newly differentiated IHCs and OHCs from the cochlear basal turn at E15.5 and E16.5, respectively (Fig. 4 and data not shown). At these stages, the actin labeling did not enable discrimination of the future stereocilia from the surrounding microvilli. By E18.5, every hair cell along the cochlea stained positive for the four Ush1 proteins, which were all concentrated at the tips of both the differentiated stereocilia and surrounding microvilli (Fig. 4). Pcdh15-cter and Cdh23-N1 stainings also extended along the entire length of the stereocilia, and they were particularly intense in the region of the hair bundle where the stereocilia are connected to the kinocilium (Fig. 4).

**Harmonin-b localization to the tips of emerging stereocilia is dependent on the presence of myosin VIIa and sans**

The simultaneous presence of four Ush1 proteins at stereocilia tips during the initial phase of hair bundle morphogenesis, in conjunction with their well-documented in vitro interactions, suggested that they can interact during this period of development. Because of the central role played by harmonin in these in vitro interactions (Fig.
The intermediate phase of stereociliary growth is defective in Ush1 mutant mice

Between P0 and P5, the three outward-most stereocilia rows of wild-type hair bundles showed differential row-specific elongation, ultimately leading to the characteristic staircase-like pattern of mature hair bundles (Fig. 6A). Strikingly, SEM analysis of Ush1 mutant hair bundles during this period of development showed that many stereocilia were of abnormal length (Fig. 6A,B). Many stereocilia of Myo7a^4626SB/4626SB mice were taller than those in wild-type mice (Fig. 6B). By contrast, in all the other Ush1 mouse mutants, many stereocilia of the small and, to a lesser extent, medium rows were shorter than expected, whereas stereocilia of the tall row were of normal height (Fig. 6A,B). In particular, in OHCs defective in harmonin, the majority of stereocilia from the small and medium rows did not elongate further from P0. At P5 they were in the process of regressing, and at P15 they had disappeared (Fig. 6A).

As row-specific elongation proceeded, the tips of small and medium stereocilia evolved from a round, oblate shape into an asymmetric, prolate shape in wild-type hair bundles (Fig. 6C). Such a morphological change is thought to result from tension forces applied, via the tip link, to the apical membrane of these stereocilia (Rzadzinska et al., 2004). In Ush1 mutant hair bundles, especially in IHCs, stereocilia of the small and medium rows virtually never had prolate tips, but rather had round, oblate tips (Fig. 6C), indicating that these putative tension forces might not develop properly in the absence of Ush1 proteins.

We therefore examined the distribution of interstereociliary links in the different Ush1 mutant hair bundles. During late embryonic and early postnatal stages, the lateral fibrous links that are initially...
Harmonin-b relocalization below stereocilia tips does not occur in cadherin 23- or protocadherin 15-defective mice

The additional phenotype that we identified in Ush1 mutant mice during the period of differential stereociliary growth suggests that the Ush1 proteins also cooperate in this hair bundle morphogenetic step. Therefore, we examined whether myosin VIIa, harmonin-b, cadherin 23a and protocadherin 15a/b still colocalized in wild-type stereocilia during this period of hair bundle morphogenesis. From P0 onwards, both the Cdh23-N1 and Pcdh15-cter stainings became progressively restricted towards the distal part of the growing stereocilia, so that only stereocilia tips remained immunoreactive from P5 onwards (Fig. 4). The Cdh23-N1 labeling, however, was barely detectable after P13, indicating that our antibody fails to detect the tip link (data not shown). Meanwhile, the distribution profiles of myosin VIIa and harmonin-b underwent a dramatic change at P0 and P1, respectively. From P0 onwards, the myosin VIIa immunoreactivity of stereocilia tips progressively decreased and a new bright Myo7a-F1 staining was detected near the base of stereocilia (Fig. 4). Remarkably, from P1 and P4 onwards, harmonin-b was no longer detected at stereocilia tips of IHCs and OHCs, respectively. Instead, it was detected below the tip of tall and medium stereocilia, in single spots facing the tips of the adjacent shorter stereocilia, and not in the small stereocilia (Fig. 4). Notably, this region corresponds to the upper attachment point of the tip link. The distribution profile of harmonin-b then remained largely unchanged until hair bundles were fully mature (data not shown).

Remarkably, we observed that the harmonin-H1b immunoreactivity remained at stereocilia tips and was not detected in the region of the tip link upper end in P5 Cdh23v2jv2j and Pcdh15v3jv3j hair bundles (Fig. 7A), suggesting that the two cadherins directly or indirectly control the harmonin-b switch. By contrast, Cdh23-N1 and Pcdh15-cter labelings were detected at stereocilia tips in harmonin-null mice (Fig. 7B), as in wild-type mice. This suggests that cadherin 23a and protocadherin 15a/b do not rely on the presence of harmonin isoforms for their apical distribution. Whether they require harmonin to form functional apical links, however, remains to be examined. In Myo7a4626SB/4626SB and sans-deficient Ush1g js/js mice, harmonin-b was again not detected in the stereocilia (Fig. 7A).

DISCUSSION

In this study, we identified common morphological anomalies of developing hair bundles in mouse models for the five USH1 genetic forms. This core Ush1 phenotype includes the fragmentation and misorientation of hair bundles from the earliest stages of their development, as well as stereocilia elongation and apical tip defects at later stages.

Hair bundle polarization has been described as a two-step process. From E15.5 in the mouse, the kinocilium migrates towards a lateral position, from the center to the periphery of the cell apical surface. Then, once stereocilia have differentiated and assembled into a bundle, a reorientation step occurs in which the hair bundle progressively reaches its final location on the cell apical surface, pointing towards the distal pole (Cotanche and Corwin, 1991; Denman-Johnson and Forge, 1999; Dabdoub et al., 2003). The role of the kinocilium in the differentiation, growth and assembly of the
closest microvilli into a polarized, V-shaped stereociliary bundle is still obscure. However, its directional migration towards the cell periphery before any hair bundle is recognizable strongly suggests that it has a leader role in the establishment of hair bundle polarity. Moreover, a lack of polarization or a mispolarization of the stereocilia bundles has recently been reported in mutant hair cells, in which the basal body remained in a central position or was mispositioned, respectively (Jones et al., 2008). In all Ush1 mouse mutants analyzed here, kinocilia were most frequently mispositioned. Nevertheless, virtually no kinocilia were found at the center or within the medial half of the hair cell apical surface, indicating that the first step of hair bundle polarization is not affected in Ush1 mutant mice. Moreover, the polarization of Vangl2 and frizzled 3, two essential components of the core PCP pathway (for a review, see Wang and Nathans, 2007) that precedes and participates in the polarized positioning of the kinocilium (Montcouquiol et al., 2003; Wang et al., 2006; Deans et al., 2007), occurred normally at the polarized positioning of the kinocilium (Montcouquiol et al., 2003; Wang and Nathans, 2007) that precedes and participates in the polarized positioning of the kinocilium. Indeed, cadherin 23a or protocadherin 15a/b either at the hair cell apical junctions or at the basal body. Instead, our findings suggest that the proper final orientation of the hair bundle requires both the cohesiveness of the stereociliary bundle and its connection to the kinocilium. Indeed, cadherin 23a and protocadherin 15a/b transmembrane isoforms were first detected in the presumptive stereocilia as soon as the kinocilium had reached the cell periphery (at around E16). This indicates that the formation of the stereokinociliary and interstereociliary links made by these cadherins precedes the hair bundle reorientation step. Consistently, interstereociliary and stereokinociliary links have been detected in a majority of hair cells along the cochlea at E17.5 (Goodyear et al., 2005). Moreover, in Cdh23 av3J/v2J and Pcdh15 av3J/v2J mutant mice that lack these isoforms, kinocilia were often dissociated from the stereociliary bundles and showed, on average, greater deviations than those of other Ush1 mutant hair cells. From our results, we can draw the general conclusion that the part of the hair bundle emerging from the apical cell surface, together with the expected cytoskeletal connection between the basal body and cell-cell junctions (Jones et al., 2008), contributes to the determination of the hair bundle final position. Incidentally, the observation that hair bundles in Cdh23 av3J/v2J and Pcdh15 av3J/v2J mutant mice often have abnormal

Fig. 6. Elongation and tip defects of stereocilia in Ush1 mutants during postnatal stages. (A) Hair bundle maturation in P0 to P15 wild-type and Ush1C−/− OHCs from the end of the apical cochlear turn. At P0, mutant OHC bundles are composed of four to six stereocilia rows almost equal in height, resembling their wild-type counterparts. From this stage onwards, wild-type stereocilia undergo differential growth, depending on the row they belong to. In the absence of harmonin, small and medium stereocilia rows show marked elongation defects from P2 onward, and most of the stereocilia from these rows have disappeared by P15. By contrast, stereocilia from the tall row are of normal length at all stages examined. Note that some small and medium row stereocilia located near clump vertices show some elongation at first (see black, green and red lines running along the tall, medium and small rows of stereocilia, respectively, in P2 and P5 wild-type and mutant OHC bundles). (B) OHC bundles of P5 Myo7a+4626SB/4626SB, Cdh23 av3J/v2J, Pcdh15 av3J/v2J and Ush1G−/− mice (view from the end of the apical cochlear turn). Note the abnormal height of many stereocilia of the medium row and the frequent absence of small stereocilia in mutant hair bundles. (C) Mid-cochlear IHC bundles of P5 wild-type, Myo7a+4626SB/4626SB, Ush1C−/−, Cdh23 av3J/v2J, Pcdh15 av3J/v2J and Ush1G−/− mice. As differential elongation occurs, stereocilia of the medium row in wild-type hair bundles acquire a particular prolate shape (see gray lines in the wild-type IHC inset), which could result from tension forces applied to their tip membrane (Rzadzinska et al., 2004). Medium stereocilia of Ush1 mutant IHCs, however, never display prolate tips (see insets). In addition, apical links (arrowheads) are either absent (Cdh23 av3J/v2J and Pcdh15 av3J/v2J), or appear sparser (Myo7a+4626SB/4626SB, Ush1C−/− and Ush1G−/−) in the Ush1 mutants, compared with the controls. Scale bars: 1 μm.
myosin IIIa (Schneider et al., 2006). However, the distribution of myosin VIIa is reminiscent of that induced by mutant forms of growth). The increased elongation of stereocilia in the absence of compared with the absence of any other Ush1 protein (decreased stereociliary elongation. At first glance, the absence of myosin VIIa phenotype of all the mutants. Supporting this proposal, in the vestibule, in which extensive epithelium remodeling by convergent extension does not occur, hair bundles of Ush1 mutants are rarely fragmented in clumps and display only minor orientation abnormalities, whereas they show significant differential growth defects (G.L., unpublished).

During the intermediate phase of hair bundle morphogenesis, the phenotype of all the Ush1 mutant mice examined includes abnormal stereociliary elongation. At first glance, the absence of myosin VIIa has an opposite effect on stereocilia growth (increased growth) compared with the absence of any other Ush1 protein (decreased growth). The increased elongation of stereocilia in the absence of myosin VIIa is reminiscent of that induced by mutant forms of myosin IIIa (Schneider et al., 2006). However, the distribution of myosin IIIa, which is restricted to stereocilia tips, indicates a local role for this protein in acting directly on the machinery controlling actin polymerization, whereas the presence of myosin VIIa along the stereocilia shafts rather suggests that it acts as a conveyer of key regulators of actin polymerization towards stereocilia tips. The stereocilia growth defect observed in the other Ush1 mutants is unprecedented. Indeed, it differs from that observed in mice deficient for myosin XVa, whirlin or espin (Mburu et al., 2003; Probst et al., 1998; Sjostrom and Anniko, 1992; Zheng et al., 2000) in that it spares stereocilia of the tallest row. The concomitant appearance of this defect and switch of the harmonin-b staining from the stereocilia tip to the upper attachment point of the tip link, in conjunction with the involvement of cadherin 23 and protocadherin 15 as tip link components (Ahmed et al., 2006; Kazmierczak et al., 2007; Siemins et al., 2004), strongly suggest that the stereocilia elongation defect of Ush1 mutants results from insufficient tension forces applied by the tip links on the tips of small and medium stereocilia. Along this line, pulling forces applied to actin filaments have been predicted to control actin polymerization (Hill and Kirschner, 1982), and a mechanism involving formins in this process has been proposed (Kozlov and Bershadsky, 2004). Finally, regarding the differential growth of the tallest stereocilia row, the observation that the length of these stereocilia is not affected when the kinocilium either lacks its axonemal part (Jones et al., 2008) or as a consequence of the loss of some interstereociliary links, as in the case of Cdh23<sup>3v2v2</sup> and Pcdh15<sup>av3av3</sup> mutants, this study), suggests that the kinocilium does not play a crucial role in this process.

Additional lines of evidence suggest that harmonin-b anchors the tip link upper end (likely to be made of cadherin 23) to the stereocilia actin core, hence participating in the transmission of the above-mentioned tension forces. Firstly, direct interactions of harmonin-b with cadherin 23 and F-actin have been shown in vitro (Adato et al., 2005; Boeda et al., 2002; Siemins et al., 2002). Secondly, the harmonin-b immunoreactivity switch does not occur in Cdh2<sup>3v2v2</sup> and Pcdh15<sup>av3av3</sup> mice that do not have any detectable tip links.
Thirdly, small and medium stereocilia have oblate-shaped tips in Cdh23−/−21v23, Pcdh15v33v31 and Ush1c−/− mutants, instead of the normal prolate-shaped tips that are believed to result from the traction force exerted by the tip link on the apical membrane (Rzadzinska et al., 2004; Prost et al., 2007). Notably, the elongation defect in sans-deficient Ush1g−/− mice might result from the absence of harmonin-b, which was never detected in the stereocilia of these mice.

In conclusion, our results on Ush1 mutant mice shed new light on the cellular mechanisms involved in hair bundle morphogenesis. In particular, they unravel the role of interstereociliary and stereokinociliary links in hair bundle cohesion and orientation at early developmental stages. Moreover, they point to a previously unrecognized role of the tip link in stereocilia differential growth, in addition to its well-established role in mechanoelectrical transduction.

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