A New Compartment at Stereocilia Tips Defined by Spatial and Temporal Patterns of Myosin IIIa Expression

Mark E. Schneider,1,2 Andréa C. Dose,2* Felipe T. Salles,1* Weise Chang,1 Floyd L. Erickson,3 Beth Burnside,2 and Bechara Kachar1

1Section on Structural Cell Biology, National Institute on Deafness and Other Communication Disorders, National Institutes of Health, Bethesda, Maryland 20892; 2Department of Molecular and Cell Biology, University of California, Berkeley, California 94720; and 3Department of Biological Sciences, Salisbury University, Salisbury, Maryland 21801

Class III myosins are motor proteins that contain an N-terminal kinase domain and a C-terminal actin-binding domain. We show that myosin IIIa, which has been implicated in nonsyndromic progressive hearing loss, is localized at stereocilia tips. Myosin IIIa progressively accumulates during stereocilia maturation in a thimble-like pattern around the stereocilia tip, distinct from the cap-like localization of myosin XVa and the shaft localization of myosin Ic. Overexpression of deletion mutants for functional domains of green fluorescent protein (GFP)–myosin IIIa shows that the motor domain, but not the actin-binding tail domain, is required for stereocilia tip localization. Deletion of the kinase domain produces stereocilia elongation and bulging of the stereocilia tips. The thimble-like localization and the influence myosin IIIa has on stereocilia shape reveal a previously unrecognized molecular compartment at the distal end of stereocilia, the site of actin polymerization as well as operation of the mechanoelectrical transduction apparatus.

Key words: hair cells; myosin IIIa; stereocilia; hearing; myosin XVa; myosin Ic

Introduction

The staircase-shaped stereocilia bundle of the inner ear sensory hair cells detect mechanical stimuli by the activation of mechanoelectrical transduction (MET) channels at the stereocilia tips (for review, see Ricci et al., 2006). Stereocilia formation, hair bundle organization, and mechanosensory function depend on several unconventional myosins that when mutated cause stereocilia disruption and deafness. For example, myosin XVa is essential for stereocilia elongation (Belyantseva et al., 2002; Rzadzinska et al., 2005), myosin VIIa is required for maintaining proper cohesiveness of the stereocilia bundle (Boeda et al., 2002; Kros et al., 2002), and myosin VI is involved in maintaining the tapered base of the stereocilia (Seiler et al., 2004). Myosin Ic is another unconventional myosin that has been implicated in adaptation of the MET response (for review, see Gillespie and Cyr, 2004).

MYO3A, one of two human class III myosins (Dose and Burnside, 2000), was shown to play a critical role in inner ear function because mutations in the motor domain of this gene lead to the progressive, nonsyndromic hearing loss DFNB30 (Walsh et al., 2002). Myosin IIIa is also expressed in the vertebrate retina (Dose et al., 2003), and its Drosophila homolog, NINAC, is expressed in photoreceptors. NINAC interacts with actin filaments and with INAD, a PDZ (postsynaptic density-95/Discs large/zona occludens-1) scaffolding protein, and has been proposed to translocate and assemble proteins of the phototransduction signaling complex (Lee and Montell, 2004; Wes et al., 1999). Class III myosins share structural features with myosins XVa, VIIa, and Ic including an N-terminal canonical motor domain and a short regulatory neck domain containing a variable number of IQ motifs (Dose and Burnside, 2000; for review, see Sellers, 2000). Myosin III proteins are unique in that they contain an N-terminal kinase domain and undergo autophosphorylation (Ng et al., 1996; Komaba et al., 2003). Deletion of the kinase domain does not prevent myosin IIIa localization to HeLa cell filopodia tips (Les Erickson et al., 2003), but this domain is essential for normal phototransduction in Drosophila (Porter and Montell, 1993). The myosin IIIa C terminus contains two conserved signature motifs termed myosin III tail homology domains, 3THDI and 3THDII (Dose et al., 2003). A point mutation in the second conserved motif, 3THDII, eliminated the actin binding that is essential for targeting a fish myosin IIIa to filopodia tips, but not to photoreceptor calycal processes (Les Erickson et al., 2003; Lin-Jones et al., 2004).

We used immunolabeling to show that myosin IIIa is localized around the stereocilia tips forming a thimble-like pattern. Expression of green fluorescent protein (GFP)-tagged myosin IIIa showed that the motor domain, but not the conserved tail actin-binding domain, is required for stereocilia tip localization. Furthermore, deletion of the unique 5’ kinase domain enhances the accumulation of myosin IIIa and results in elongation of the stereocilia and bulging of the stereocilia tips. We argue that the thimble-like...
localization and the influence myosin IIIa has on stereocilia shape reveal a previously unrecognized molecular compartment at the distal end of stereocilia, the site of actin polymerization as well as operation of the MET.

Materials and Methods

**Antibodies, expression plasmids, and in situ probes.** An affinity-purified polyclonal antibody (PB638) was developed in a rabbit immunized with a synthetic peptide (NYPDYRRLLSQRQR) matching the C-terminus sequence of mouse myosin IIIa (Princeton Biomolecules, Langhorne, PA). Two other affinity-purified antibodies, 5525 and 5526, were generated previously (Dose et al., 2003) in two rabbits both given injections of a peptide corresponding to the last 22 aa of the bass (*Morone saxatilis*) myosin IIIa C terminus. The epitope used for the bass antibody production contained extensive sequence homology to the mouse, rat, and human myosin IIIa tail end (Dose et al., 2003). The specificity of all three myosin IIIa antibodies was tested using ectopic expression of GFP-tagged plasmid DNA for the human and bass myosin IIIa in COS-7 cells (data not shown). Affinity-purified rabbit polyclonal antibodies and GFP-tagged plasmids to myosin XVA (Anderson et al., 2000) and myosin Ic (Dumont et al., 2002) were obtained from T. Friedman [National Institute on Deafness and Other Communication Disorders (NIDCD), National Institutes of Health (NIH), Bethesda, MD] and P. Gillespie (Vollum Institute, Portland, OR), respectively. Monoclonal anti-β-tubulin (clone D66) was obtained from Sigma (St. Louis, MO).

We characterized the phenotypes of cells transfected with bass and human myosin IIIa and mutations of specific domains subcloned into pEGFP-C1, -C2, -C3 (GFP) vectors (Invitrogen, Carlsbad, CA). Bar diagrams of the functional domains of GFP-tagged constructs and their abbreviations used are shown in Figure 5, a and b. The bass constructs used were as follows: WT, wild-type bass GFP–myosin IIIa (amino acids 1–1838); MD, motor dead bass GFP–myosin IIIa (with an R-to-A mutation at amino acid position 507); ΔK, bass GFP–myosin IIIa Δ kinase (amino acids 361–1838); ΔT3HDI, bass GFP–myosin IIIa (amino acids 1–1816). The human constructs used were as follows: WT, human GFP–myosin IIIa (amino acids 1–1615); ΔK, human GFP–myosin IIIa Δ kinase (amino acids 340–1615); Δ34, human GFP–myosin IIIa Δ exon 34 (deletion of exon 34, which results in missing C-terminal two-thirds of 3T3HDI, but 3T3HDI is still in frame); ΔK–34, human GFP–myosin IIIa Δ kinase, Δ exon 34 (deletion of both domains); Δ34–33, human GFP–myosin IIIa, Δ exon 33 and 34 (out of frame after first five amino acids of 3T3HDI and therefore missing 3T3HDI as well); ΔK–33–34, human GFP–myosin IIIa Δ kinase, Δ exons 33 and 34 (deletion of all three domains).

**In situ hybridization for myosin IIIa.** Freshly dissected inner ear tissues harvested from 8–d-old rats were fixed in a mixture of 4% PFA and 0.001 U/ml Alexa Fluor 568 phallolidin (Invitrogen), washed with PBS, and mounted using Prolong anti-fade (Invitrogen). Some samples were processed for antigen retrieval. Briefly, fixed and permeabilized samples were transferred to 0.1 m citrate buffer, pH 7.0, and heated to 60°C for 30 min before blocking overnight and processed as before. Fluorescence confocal images were obtained with a Nikon (Tokyo, Japan) microscope using a 60× or 100× 1.4 numerical aperture (NA) objective and a spinning wheel confocal unit (PerkinElmer, Wellesley, MA). We analyzed at least three samples from three animals at each age used, and representative images are shown.

**Tissue preparation for electron microscopy.** Freshly dissected inner ear tissues harvested from 8–d-old rats were fixed in a mixture of 4% PFA and 0.001 U/ml Alexa Fluor 568 phallolidin, glycinated, plunged into liquid Freon 22, and promptly transferred to liquid nitrogen. Frozen samples were freeze substituted in 1.5% uranyl acetate in absolute methanol at −90°C, infiltrated with Locrycel HM-20 resin at −45°C, polymerized with UV light, thin-sectioned, and immunogold labeled as described previously (Dumont et al., 2001). Stained sections were viewed and photographed with a Leo 922 transmission electron microscope equipped with an energy filter for contrast tuning.

**Cultures and transfection of rat inner ear tissue.** After CO2 anesthesia, postnatal day 0 (P0) to P4 rat pups were killed in accordance with NIH guidelines, and their temporal bones were isolated and placed in L-15 medium (Invitrogen). Pieces of organ of Corti and maculae of the utricle, saccule, and the ampullae were dissected and attached to coverslips previously coated with 150 µg/ml Cell Tak (BD Biosciences, San Jose, CA) as described previously (Rzadzinska et al., 2004). Cultures were maintained in DMEM (Invitrogen) supplemented with 5–7% fetal bovine serum (FBS) (Invitrogen) and 1.5 µg/ml ampicillin and kept at 37°C and 5% CO2. Plasmid DNAs encoding the various GFP–myosin IIIa constructs were precipitated onto 1 µm gold particles at 50 µg of DNA to 25 mg of 1 µm gold particles and loaded into cartridges following the Helios Gene Gun protocol (Bio-Rad, Hercules, CA). Tissue explants were transfected with these gold carriers using the Helios Gene Gun using 85–95 psi helium pressure and incubated for up to 48 h. Samples were washed briefly with PBS, fixed at various times in 4% PFA for at least 1 h, and permeabilized with 0.5% Triton X-100 in PBS containing Alexa Fluor 568 phallolidin for 30 min before washing and mounting on a microscope slide with the anti-fade. The efficiency of transfection ranged from 0 to 3 hair cells per tissue explant, and we examined at least 20 transfected cells for each construct used.

**Culture and transfection of COS-7 cells.** Simian fibroblast COS-7 cells were plated on coverslips and maintained at 37°C in DMEM supplemented with 10% FBS. Cultures were transfected using the GeneJuice system (Novagen, San Diego, CA) according to the manufacturer’s directions. Cells were fixed at various times after transfection for 20 min in 4% PFA in PBS, permeabilized for 10 min in 0.5% Triton X-100 in PBS, processed for immunocytochemistry or counterstained with Alexa Fluor 568 phallolidin for 20 min, and mounted on a microscope slide with the anti-fade. Live-cell imaging of transfected COS-7 cells was performed in L-15 medium. Time-lapse confocal Z-stacks were collected using a Nikon Eclipse microscope and PerkinElmer Ultraview confocal with a 100× 1.4 NA objective. The extended focus rendering was performed using Velocity 3.7.0 (Improvement, Coventry, UK). We examined >40 cells over periods of 20 min each.

In situ hybridization experiments were performed as described previously (Wu and Oh, 1996). Briefly, tissues were fixed in 4% PFA in PBS overnight in the cold room and processed through graded sucrose solutions (3–30%) for cryoprotection before cryosectioning. Digoxigenin-labeled riboprobes were produced using DNA templates as described above. The results of the in situ hybridization were then visualized using nitroblue tetrazolium/5-bromo-4-chloro-3-indolyl phosphate chromogenic development procedure.

**Results**

**Myosin IIIa localized to tips of developing stereocilia of inner ear hair cells**

We used three affinity-purified myosin IIIa-specific antibodies (PB638, 5525, and 5526) to characterize the steady-state distribution and temporal patterns of expression of myosin IIIa in developing and mature hair bundles from P0 through adult (>P30)
Myosin IIIa expression increased as stereocilia elongated in the cochlea

We observed the highest levels of myosin IIIa immunoreactivity at tips of stereocilia between P6 (Fig. 2a,b) and P10 (Fig. 2c,d) in the rat cochlea. Because stereocilia range between 200 and 500 nm in diameter, the fluorescence images are diffraction limited. However, in the longest and widest stereocilia of the cochlea (from the inner hair cells of the apical turn), the tips exhibited a clearly distinguished thimble-like pattern of fluorescence. Confocal images of side views (Fig. 2a,d) or of top views (Fig. 2e–g) of the tallest row of stereocilia showed a ring pattern of labeling surrounding the unlabeled stereocilia actin core. The staining pattern for the middle and short rows of stereocilia often showed an asymmetric pattern with labeling more concentrated toward the tallest row of stereocilia (Fig. 2d, arrows). The pointed fluorescence pattern was consistent with the asymmetric pointed shape of the second row of stereocilia as seen by electron microscopy (Kachar et al., 2000). In the outer hair cells, the middle-row stereocilia often showed an apparent higher level of myosin IIIa immunoreactivity (Figs. 1e, 2b).

Changes in myosin IIIa immunoreactivity during late stages of stereocilia maturation

Using our standard protocol for immunofluorescence, we initially observed that myosin IIIa immunofluorescence levels dropped from the peak of expression between P6 and P10 (Fig. 2a–d) to very low levels in the adult cochlea (Fig. 3a–f), whereas the thimble-pattern of fluorescence appeared more restricted to the stereocilia tip (compare Figs. 2a, 3a,g). In the hair cells of the basal turn of adult guinea pig cochlea, myosin IIIa staining was barely detectable at the stereocilia tips (Fig. 3f). No labeling was detected in the rat or mouse adult hair cell stereocilia (Fig. 3d,e). In vestibular hair cells, myosin IIIa was localized at the tips of stereocilia and the thimble-like banding pattern could be recognized up to P20 (Fig. 3j). In stereocilia of adult vestibular hair cells, the fluorescence intensity of the labeling decreased and appeared as a series of puncta (Fig. 3k).

The progressive loss of immunofluorescence intensity at the later stage of maturation of the stereocilia observed in our immunofluorescence assay was inconsistent with the reported involvement of MYO3A in late-onset deafness (Walsh et al., 2002). We explored the possibility that the loss of immunoreactivity might be caused by some form of epitope masking. We used an antigen retrieval step, which consisted of incubation of the fixed tissue for 30 min at 60°C in citrate buffer, and observed a robust recovery of the immunolabeling signal intensity in the stereocilia of all adult hair cells, including adult rat, mouse, and guinea pig cochlear (Fig. 3g–i) and rat vestibular (Fig. 3i) hair cells. With this antigen retrieval protocol (data not shown), we also confirmed the progressive increase of immunofluorescence intensity and labeling pattern observed during the early developmental stages with our cochlea. Immunofluorescence experiments were performed using all three anti-myosin IIIa affinity-purified antibodies in rat and mouse inner ear sensory epithelia. The overall results were indistinguishable regardless of the antibodies and rodent species used, and we show in the figures representative results using PB638 in rat preparations. In the basal turns of the rat cochlea, stereocilia emerge from the apical surface of the inner and outer hair cells at embryonic day18, and by P21, the hair bundles have reached their full, mature length and lost their kinocilium (Tilney et al., 1988; Kaltenbach et al., 1994; Zine and Romand, 1996). Maturation of inner hair cells of the basal region precedes the maturation of outer hair cells, and development proceeds toward the apex of the cochlea over a few days (Zine and Romand, 1996). In the comparitively immature outer hair cells of the apical regions of the P0 rat cochlea, no myosin IIIa immunolocalization could be detected at the tips of the emerging stereocilia (Fig. 1a). In the longer stereocilia of the inner hair cells of the basal turn of the P0 cochlea and in the vestibular hair cells where the hair bundle already has the characteristic staircase shape, we observed distinct myosin IIIa immunolabeling at the tips of stereocilia (Fig. 1b,c). By P3, myosin IIIa labeling was observed at stereocilia tips of both inner and outer hair cells of the apical turns of the cochlea (Fig. 1d,e). Myosin IIIa immunolabeling could not be detected in the kinocilium, which is present in all vestibular hair cells and in the immature cochlear hair cells (Fig. 1c–e).
standard protocol and the late confinement or restriction of the thimble-like fluorescence within the stereocilia tips.

We sought to independently confirm the myosin IIIa expression throughout hair cell development and maturation using in situ hybridization. We hybridized mouse cryosections of the cochlea and ampullae from P0, P8, P15, P30, and P50 with a 1 kb riboprobe directed to the mouse myosin IIIa N terminus and used a 1 kb myosin XVa probe as a hair cell marker (Liang et al., 1999). We observed a strong hybridization signal from inner and outer hair cells at P8 (supplemental Fig. 1a, available at www.jneurosci.org as supplemental material) as well as from hair cells of the ampullae (supplemental Fig. 1b, available at www.jneurosci.org as supplemental material). Moreover, clear hybridization labeling in the P50 mouse cochlear and vestibular hair cell (supplemental Fig. 1c,d, available at www.jneurosci.org as supplemental material) was observed. Positive hybridization experiments in the adult cochlea suggested that myosin IIIa is expressed in adult hair cells, consistent with our immunocytochemical observations.

We extended our findings on myosin IIIa distribution by studying bullfrog hair cells with our standard protocol and with the added antigen retrieval step. The thimble-like distribution of myosin IIIa in stereocilia was observed under either protocol in this non-mammalian vertebrate (Fig. 4). In the frog saccule and utricle hair cells, the myosin IIIa immunofluorescence of the middle-row stereocilia exhibited a pointed pattern (Fig. 4a, arrow, b). In the very long stereocilia of the ampullae hair cells, the myosin IIIa immunofluorescence extended from the tips downward along the stereocilia shaft in a comet-like pattern (Fig. 4d–f). Some of the longest stereocilia of the ampullae exhibited slightly bulbous or club-shaped tips, and these unusual phenotypes were also strongly positive for myosin IIIa labeling, especially over the enlarged region of the tips (Fig. 4c–f, arrowheads). We observed these various degrees of bulging of the tips in ~1–2% of the cells.

We also evaluated myosin IIIa localization by postembedding immunogold electron microscopy of rat (supplemental Fig. 2a–c, available at www.jneurosci.org as supplemental material) and chick (supplemental Fig. 2d, available at www.jneurosci.org as supplemental material) hair cells. Labeling could be observed around the stereocilia tips in longitudinal and perpendicular sections in a pattern consistent with the thimble-like staining pattern of the immunofluorescence confocal images.

**Figure 2.** Myosin IIIa immunoreactivity at stereocilia tips peaks between P6 and P10, forming a thimble-like pattern. Confocal images of organ of Corti hair cell stereocilia from P6 (a, b), and P10 (c–g) show myosin IIIa immunofluorescence (green) at the tips of stereocilia counterstained in red for actin. a, d, A thimble-like pattern can be observed in the larger stereocilia of the inner hair cells. b, In the outer hair cells, the second row of stereocilia in each bundle often show higher levels of immunoreactivity than both their taller and shorter neighbors. c, d, In the inner hair cells, the immunofluorescence at the second-row stereocilia often exhibits a pointed shape (arrows). e–g, A confocal optical section across the tips of stereocilia of the inner hair cells (e; close up views in f and g) shows the thimble-like pattern of immunolabeling of myosin IIIa around the tips of stereocilia. Scale bars: a–c, 5 μm; d, f, g, 2 μm.

**GFP–myosin IIIa was targeted to the tips of stereocilia in transfected hair cells**

To assess the ability of hair cells to target and properly compartmentalize exogenous myosin IIIa to the tips of stereocilia, we transfected hair cells of P4–P11 rat inner ear sensory epithelia cultures with bass (Fig. 5a) or human (Fig. 5b) GFP-tagged myosin IIIa. Within 18–48 h after hair cell transfection, myosin IIIa was concentrated at the tip of each stereocilium in transfected cells of organ of Corti and vestibular epithelia (Fig. 5c–f). High-magnification of the well resolved stereocilia images suggest that GFP–myosin IIIa forms a thimble-like pattern at the stereocilia tips (Fig. 5d,f, arrowheads) similar to that observed by myosin IIIa immunolocalization. We compared transfected hair cells from the organ of Corti with nontransfected neighboring cells to assess the effect of GFP–myosin IIIa overexpression on promoting actin elongation and stereocilia lengthening. Unlike our observations with GFP–myosin XVa (H. W. Lin and B. Kachar, unpublished data), GFP–myosin IIIa overexpression did not produce evident changes in stereocilia length (data not shown).

**Targeting and structural effects of mutated myosin IIIa in stereocilia**

Previous studies on myosin IIIa have identified critical domains in which mutations affected the localization pattern of heterologously expressed GFP–myosin IIIa in filopodia of cultured cell
and bulging were evident after visual inspection of the fluores-
cence images (in the >40 transfected cells observed) but were
difficult to quantify because the entire stereocilia bundle staircase organization was often disrupted (Fig. 5h–j and supplemental
Fig. 3a,c, available at www.jneurosci.org as supplemental material). Interestingly, the ΔK-transfected nonsensory cells of either the organ of Corti or vestibular sensory epithelia, which commonly have short microvilli on their surface, exhibited unusually long and irregular actin protrusions (supplemental Fig. 3b, available at www.jneurosci.org as supplemental material). These long actin protrusions were not observed when these cells were transfected with the full-length GFP–myosin IIIa (supplemental Fig. 3b, inset, available at www.jneurosci.org as supplemental material).

A conserved tail domain motif, 3TH-DII, has been shown previously to bind actin and was required for localization of GFP–myosin IIIa to filopodia tips in a heterologous expression system (Les Erickson et al., 2003). We tested whether elimin-
ation of the 3THDII region from either the bass or human transcripts affected the targeting of myosin IIIa and influenced the shape of the stereocilia. We tested two mut-
ant variants of myosin IIIa associated with the 3THD domains. The human GFP–myosin IIIa Δ33–34 lacks exons 33 and 34, which removes all but the first five amino acids of 3THDII and also creates a frame shift altering the rest of the tail, thereby eliminating 3THDII. The myosin IIIa Δ34 variant removes the second two-thirds of 3THDII, because it does not cause a frame shift, so 3THDII is still present at the C terminus. The Δ34 is a naturally occurring splice variant seen in mouse (Walsh et al., 2002), and the Δ33–34 variant was identi-
fied among other variants in human cDNA (A. C. Dosé, unpublished results). Both of the human GFP–myosin IIIa tail variants efficiently localized to stereocilia tips and showed a phenotype indistinguishable from the wild-type myosin IIIa overexpression (Fig. 5k,l). A species difference between bass and human was observed in the localization of tail mutants in hair cells. Bass GFP–myosin IIIa missing exclusively the final 22 amino acids (3THDII) resulted in an attenuated expression in stereocilia tips (data not shown).

Because the absence of the kinase domain affected the func-
tional activity of myosin IIIa, but absence of 3THDII did not (unlike in cultured cell lines), we sought to examine the effect of mutations in both domains simultaneously. Expressed myosin IIIa lacking both the kinase domain and residues encoded by exons 33 and 34 (ΔKΔ33–34) no longer exhibited the aberrant ΔK phenotype and was indistinguishable from the wild type (Fig. 5m). However, expression of myosin IIIa lacking the kinase do-
maintaining of 3THDII, caused increased fluorescence, stereocilia elongation, and bulg-
ing at the tips (Fig. 5n and supplemental Fig. 3c, available at www.jneurosci.org as supplemental material), resembling the ΔK

difficult to quantify because the entire stereocilia bundle staircase organization was often disrupted (Fig. 5h–j and supplemental
Fig. 3a,c, available at www.jneurosci.org as supplemental material). Interestingly, the ΔK-transfected nonsensory cells of either the organ of Corti or vestibular sensory epithelia, which commonly have short microvilli on their surface, exhibited unusually long and irregular actin protrusions (supplemental Fig. 3b, available at www.jneurosci.org as supplemental material). These long actin protrusions were not observed when these cells were transfected with the full-length GFP–myosin IIIa (supplemental Fig. 3b, inset, available at www.jneurosci.org as supplemental material).

A conserved tail domain motif, 3TH-DII, has been shown previously to bind actin and was required for localization of GFP–myosin IIIa to filopodia tips in a heterologous expression system (Les Erickson et al., 2003). We tested whether elimin-
ation of the 3THDII region from either the bass or human transcripts affected the targeting of myosin IIIa and influenced the shape of the stereocilia. We tested two mut-
ant variants of myosin IIIa associated with the 3THD domains. The human GFP–myosin IIIa Δ33–34 lacks exons 33 and 34, which removes all but the first five amino acids of 3THDII and also creates a frame shift altering the rest of the tail, thereby eliminating 3THDII. The myosin IIIa Δ34 variant removes the second two-thirds of 3THDII, because it does not cause a frame shift, so 3THDII is still present at the C terminus. The Δ34 is a naturally occurring splice variant seen in mouse (Walsh et al., 2002), and the Δ33–34 variant was identi-
fied among other variants in human cDNA (A. C. Dosé, unpublished results). Both of the human GFP–myosin IIIa tail variants efficiently localized to stereocilia tips and showed a phenotype indistinguishable from the wild-type myosin IIIa overexpression (Fig. 5k,l). A species difference between bass and human was observed in the localization of tail mutants in hair cells. Bass GFP–myosin IIIa missing exclusively the final 22 amino acids (3THDII) resulted in an attenuated expression in stereocilia tips (data not shown).

Because the absence of the kinase domain affected the func-
tional activity of myosin IIIa, but absence of 3THDII did not (unlike in cultured cell lines), we sought to examine the effect of mutations in both domains simultaneously. Expressed myosin IIIa lacking both the kinase domain and residues encoded by exons 33 and 34 (ΔKΔ33–34) no longer exhibited the aberrant ΔK phenotype and was indistinguishable from the wild type (Fig. 5m). However, expression of myosin IIIa lacking the kinase do-
maintaining of 3THDII, caused increased fluorescence, stereocilia elongation, and bulg-
ing at the tips (Fig. 5n and supplemental Fig. 3c, available at www.jneurosci.org as supplemental material), resembling the ΔK

Figure 3. Myosin IIIa epitopes are masked as stereocilia mature. The immunoreactivity to myosin IIIa (green) changes at the late stages of stereocilia development (counterstained for actin in red). a–c, Inner hair cells from the apical turn of the (rat) cochlea from P16 (a), P18 (b), and P24 (c) showing decreased immunofluorescence and condensation into the stereocilia tip. d, f, Fluorescence is absent in the tips of stereocilia of inner hair cells of the basal turn of the adult rat (d) and mouse (e) cochlea and barely detectable in the adult guinea pig (f). g–i, After antigen retrieval, myosin IIIa immunofluorescence is observed at the tips of stereocilia in the inner hair cells of the basal turn of the adult rat (g), mouse (h), and guinea pig (i) cochlea. j–l, Myosin IIIa immunofluorescence at the tips of stereocilia of P20 (j) and adult (k, l) rat vestibular hair cells without (j, k) and with (l) the antigen retrieval step. Scale bars: a–i, 5 μm; j, k, 2 μm.
phenotype. Similarly, supporting cells expressing ΔK-34 showed elongated and irregular microvilli with GFP fluorescence at their tips. These data suggest that the presence of 3THDII is required for the ΔK phenotype.

Targeting and effect of mutants of myosin IIIa in COS-7 cells
We also evaluated the distribution and effects of GFP–myosin IIIa and the various myosin IIIa mutants on filopodia in transfected COS-7 cells (Fig. 6). Under our culture conditions, COS-7 cells exhibited very few filopodia. When COS-7 cells were transfected with GFP–myosin IIIa, the fluorescence signal was observed at the tips and along the very few filopodia even after 60 h of transfection (Fig. 6a, arrows). On the other hand, GFP–myosin IIIa ΔK caused a progressive increase in the number of filopodia that could be clearly visualized in all transfected cells after 24 h (Fig. 6b) and 48 h (Fig. 6c). Myosin IIIa mutants MD, Δ34, Δ33–34, and ΔK-33–34 all failed to localize to filopodia (Fig. 6d–g). When prepared as a construct with just the 3THDII region but missing most of 3THD1 (ΔK-34), the kinase mutation increased the number of myosin IIIa-positive filopodia in COS-7 cells (Fig. 6h, arrows). The number of filopodia seen in ΔK-34 transfected cells was less than that induced by the kinase domain deletion (ΔK) alone (Fig. 6, compare h, i). Myosin IIIa ΔK accumulated at the distal ends of filopodia, and often the fluorescence extended toward the base of the filopodia, forming a comet-like pattern (Fig. 6i, arrowhead). We examined the dynamics of GFP–myosin IIIa ΔK fluorescence in filopodia of COS-7 cells in time-lapse video recordings after 24 h after transfection. Surprisingly, although the numerous filopodia exhibited active pivoting motions about their base (supplemental movie 1, available at www.jneurosci.org as supplemental material) and intrafilopodial movements of GFP–myosin IIIa fluorescence puncta (supplemental movie 2, available at www.jneurosci.org as supplemental material), filopodial extension and retraction (Faix and Rottner, 2006) were rarely detected within the 20 min time-lapse recordings.

Myosin IIIa localization in stereocilia is distinct from myosin Ic and myosin XVa
To compare the localization of myosin IIIa with other stereocilia myosins reported to be present at the stereocilia tip, we examined in comparable experiments the precise compartmentalization of three unconventional

Figure 4. Myosin IIIa localization at regular and club-shaped stereocilia tips in the bullfrog inner ear. Immunofluorescence confocal images of utricle (a, b) and ampullae (c–f) show myosin IIIa immunoreactivity around the tips of stereocilia of adult frog vestibular hair cells. a. The arrow shows the thimble-like and pointed shape of the labeling in some middle-row stereocilia. b. The same pattern of labeling is observed after antigen retrieval. c–f. Some of the very long stereocilia of the ampullae show higher levels of immunoreactivity and bulged or club-shaped tips. In some stereocilia (d–f), the immunofluorescence may extend down the shaft for several micrometers. Scale bars, 2 μm.

Figure 5. Overexpression of myosin IIIa functional mutants in hair cells. a, b. Schematic representation of bass (a) and human (b) GFP–myosin IIIa mutants. c–n. Confocal images of cultured organ of Corti and vestibular hair cells transfected for 18–24 h with GFP–myosin IIIa mutants (green) and counterstained for actin with phalloidin (red). c–f. Human (c, d) or bass (d, f) wild-type GFP–myosin IIIa localizes at the tips of stereocilia of transfected inner hair cells (c, d) or vestibular hair cells (e, f). g, Inactivation of the bass myosin IIIa motor domain causes GFP–myosin IIIa to accumulate in the cytoplasm and confirms that stereocilia tip localization requires a functional motor domain. h–j. Images of accumulation of fluorescence and extension and bulging of the tips of stereocilia in auditory (h) and vestibular (i, j) hair cells transfected with either bass GFP–myosin IIIa ΔK (h) or human GFP–myosin IIIa ΔK (i, j). k, l. The deletion mutants GFP–myosin IIIa Δ34 (k) and GFP–myosin IIIa Δ33–34 (l) do not affect localization or the extent of accumulation in stereocilia tips. m, n. The kinase domain mutation increases the expression of myosin IIIa containing deletions of exon 34 (ΔK-34) (n) but not of a larger region of the conserved tail (ΔK-33–34) (m). Stereocilia of hair cells with ΔK-34 mutations elongate and have bulbous tips. Scale bars, 2 μm. MD, Inactive motor domain; ΔK, kinase deletion; Δ34 (human), lacking exon 34; but with 3THDII intact; Δ3TDII (bass) and Δ33–34 (human) conserved actin-binding domain 3THDII deletion or out of frame; ΔK-34 and ΔK-33–34 are combinations of deletions removing the kinase domain with deletions in the tail domain; WT, wild type.
myosins with immunofluorescence and exogenous gene transfection. We confirmed that myosin XVa immunofluorescence (Fig. 7a) was matched by the localization of GFP–myosin XVa (Fig. 7b) into a cap-like structure at the very tips of stereocilia (Rzadzinska et al., 2004). In contrast, myosin Ic was expressed in a separate and distinct compartment from myosin IIIa or XVa. Myosin Ic immunofluorescence was distributed along the entire length of the stereocilia shafts and excluded from a cap region at the tips of stereocilia (Fig. 7c, arrowheads). Overexpression of myosin Ic confirmed that myosin Ic is distributed along the entire length of the stereocilia and excluded from the stereocilia cap (Fig. 7d, arrowheads). The immunofluorescence and localization of GFP-tagged myosins XVa and Ic clearly contrasts with the localization of myosin IIIa as shown in comparable close-up views (Fig. 7e,f).

Discussion
The unique pattern of localization of myosin IIIa in stereocilia
We show that myosin IIIa forms a thimble-like pattern at stereocilia tips. In electron micrographs, the tips of stereocilia show an electron-dense structure called tip density (Kachar et al., 2000). The tip density overlaps the actin core and underlies the stereocilia membrane at the lower insertion site of the tip link. In contrast to previous reports on the distribution of myosin Ic (Garcia et al., 1998; Stegger et al., 1998), we show that it is excluded from the tip density region and distributed along the stereocilia shaft. Although myosin XVa forms a cap-like pattern at the top of the actin core matching the tip density, myosin IIIa forms a thimble-like pattern (Fig. 7e) that encircles the tip density region and extends further down the shaft of the stereocilia. Our results suggest that

---

**Figure 6.** Targeting and structural effects of mutated myosin IIIa in COS-7 cells. COS-7 cells were transfected for 24–48 h with GFP-tagged myosin IIIa constructs (green) described in Figure 5 and counterstained for actin with phalloidin (red). a, Wild-type GFP–myosin IIIa (green) is localized at the tip and along the rare filopodia (arrows). b, c, Cells transfected with ΔK for 24 h (b) and 48 h (c) show progressive accumulation of filopodia with strong fluorescence at their tips (b, arrow). b, i, Fluorescence frequently extends down the length of the filopodia (arrowheads). d–g, Mutations in the myosin motor domain (MD; d), the 3DTHII domain (Δ34; e), both conserved domains (Δ33–34; f) or the combination of deletion of the kinase domain and the tail region (ΔK-33–34; g) have few filopodia, and the myosin does not concentrate in the filopodia tips. h, i, Moderate numbers of filopodia positive for myosin IIIa at their tips are observed with the ΔK-34 mutation (compare h, i). Scale bars: a, b, d–i, 1 μm; c, 2 μm.
these three myosins have an overall distinct pattern of distribution, but, at the same time, there is some overlap of localization around the stereocilia tip (illustrated in supplemental Fig. 4, available at www.jneurosci.org as supplemental material). The thimble-like pattern of localization of myosin IIIa at stereocilia tips suggests an additional “compartment” to the structurally defined actin core and tip density. However, we cannot rule out that the myosin IIIa molecules overlying the paracrystalline actin core are part of the tip density. The potential roles of myosin IIIa at stereocilia tips are considered.

**Role of myosin IIIa in maintenance of stereocilia shape**

Although stereocilia, microvilli, and filopodia share numerous features, one key difference pertains to the stability of each protrusion. Microvilli and filopodia are formed and degraded with specific half-lives, whereas stereocilia are formed once in development, continually renewed, and last a lifetime barring an injury. Over time, stereocilia are maintained by continuous turnover, and their steady-state length and tip shape are dynamically maintained (for review, see Lin et al., 2005). Stereocilia start their elongation process before detectable amounts of the myosin IIIa are present at their tips (Fig. 1a). The progressive accumulation and subsequent condensation of myosin IIIa at the tips could contribute to stabilization of the stereocilia length and to shaping the stereocilia tip. An indication that myosin IIIa at the stereocilia tip can influence its shape comes from the result of overexpression of the kinase deletion mutants that, after accumulation of the protein at the tips, produce elongation of the stereocilia, loss of apparent rigidity, and bulging of the tip. Interestingly, high levels of myosin IIIa correlates with bulging of the tips that are occasionally observed in club-shape stereocilia in the ampullae of the frog (Fig. 4c–f).

Similar effects were seen in filopodia in cultured COS-7 (Fig. 6) and HeLa (Les Erickson et al., 2003) cell lines, calycal processes in Xenopus photoreceptors (Lin-Jones et al., 2004), and stereocilia (Fig. 5). One is tempted to hypothesize that myosin IIIa has a role in the actin bundle maintenance and stability and that the kinase-negative mutant is a hypermorphic mutation that exaggerates the myosin IIIa function.

**How does myosin IIIa reach the tips of stereocilia and how is the process regulated?**

Mutation in the myosin IIIa motor domain prevented the protein from reaching the stereocilia tips. Because actin filaments in stereocilia are uniformly polarized with their barbed end pointing to the tips, one can infer that myosin IIIa is a barbed end-directed motor and that the targeting of myosin IIIa to the tips of stereocilia depends on a motor-driven translocation along the actin filaments. The lack of the functional motor domain also prevents myosin IIIa from localizing properly in Xenopus photoreceptors (Lin-Jones et al., 2004). Interestingly, the loss of the tail actin binding site, 3THDII, prevents the myosin IIIa from localizing to the tips of filopodia as was also observed for HeLa cells (Les Erickson et al., 2003). However, in hair cells, a lack of 3THDII did not prevent accumulation of myosin IIIa at the tips of stereocilia. Similarly, myosin IIIa missing 3THDII localizes normally in Xenopus photoreceptors (Lin-Jones et al., 2004). It is possible that in these cell types the endogenous myosin IIIa is assisting in the proper localization of the mutant myosin either by a cargo-mediated interaction or even forming heterologous dimers. A myosin IIIa phenotype associated with loss of the kinase domain is more universal. In COS-7 cells, the localization to filopodial tips is more pronounced because it was also previously shown for HeLa cells (Les Erickson et al., 2003). In both hair cells and photoreceptors, lack of the kinase domain results in an apparent upregulation of activity. Interestingly, this upregulation also re-
quires 3THDII, because in hair cells, the ΔK-33–34 (lacking 3THDII) resembles the wild type and not the hypermorphic ΔK phenotype. It appears that the regulation is complex and will require more detailed mutational analysis to identify the combined role of the kinase and tail domain.

Other possible functions for myosin IIIa in stereocilia

Although some stereocilia membrane proteins can diffuse laterally to distribute along the stereocilia membrane (Grati et al., 2006), other proteins including components of the tip link complex may require active transport. Myosin XVa has been shown to shuttle the PDZ protein whirlin in stereocilia and to undergo intrafilopodial motility in transfected cultured cells (Belyantseva et al., 2005). Myosin X, an unconventional myosin similar to myosin XVa and IIIa that localizes to the tip of filopodia, was also shown to exhibit intrafilopodial motility and could shuttle filopodia components toward the tips (Berg and Cheney, 2002; for review, see Sousa and Cheney, 2005). Myosin IIIa is also in a position to shuttle cargo to the stereocilia tip. Intrafilopodial movements of GFP–myosin IIIa can be seen in COS-7 cells. Similarly, in Drosophila photoreceptors, NINAC (myosin IIIa) interacts with the adapter protein INAD, which coordinates a signaling complex via its multiple PDZ domains. INAD binds directly to rhodopsin, phospholipase C, protein kinase C, and the light-sensitive channel subunits TRP and TRPL, necessary for visual transduction (Cronin et al., 2004; Lee and Montell, 2004; Li and Montell, 2000). A mammalian INAD (Philipp and Flockerzi, 1997) was identified by reverse transcription-PCR in the cochlea (Walsh et al., 2002), and the possibility exists that myosin IIIa contributes to transport and assembly of the tip link complex, including a TRP family member that has been postulated as a likely candidate for the MET channel (for review, see Gillespie et al., 2005).

Another possibility is that myosin IIIa contributes to MET adaptation

Current models of MET adaptation suggest that myosin Ic present at either or both ends of the tip link would mediate tension on the MET gating element (Stauffer et al., 2005). Our data suggest that myosin Ic is localized along the stereocilia length but not at stereocilia tips, the site of the lower insertion of the tip link. If myosin Ic is concentrated into an adaptation complex at the upper insertion site of the tip link, then we are unable to visualize this complex or it is indistinguishable from nonadaptation myosin Ic complexes along the stereocilia (Fig. 7c). Additional myosins may play a role in MET adaptation (Kros et al., 2002). Our data do not show localization of myosin IIIa at the upper insertion site of the tip link. However, myosin IIIa is present at the lower insertion site. The higher level of immunoreactivity for myosin IIIa and the pointed shape of the labeling observed for the middle rows of stereocilia match the shape of the tip link lower insertion site at the tips of these stereocilia (Kachar et al., 2000). By forming a ring around the tip of stereocilia, myosin IIIa could also adjust tension or “tenting” of the membrane (Kachar et al., 2000) and indirectly influence tip link tension or channel gating.

What is the underlying mechanism for the late-onset deafness caused by myosin IIIa mutations?

Mutations of human MYO3A produce late-onset deafness (DFNB30) commencing at the second decade (Walsh et al., 2002), well after the onset of full hearing and after the period of stereocilia elongation. The spatial and temporal patterns of myosin IIIa localization at stereocilia tips and the effects of deletion mutants of myosin IIIa on stereocilia and other actin protrusions suggest the physiopathology of DFNB30 is likely to involve disruption of stereocilia structure and function. We cannot explain why DFNB30 patients with recessive null mutations of myosin IIIa have normal vestibular function and the hearing loss only appears decades after birth. Perhaps myosin IIIb, a short isoform of myosin III lacking 3THDII, can compensate for myosin IIIa (Dosé and Burnside, 2002). Our results showing that mutations of the distal region of the myosin IIIa tails did not reduce its ability to localize at stereocilia tips suggest that myosin IIIb may well compensate for loss of function of myosin IIIa. The localization and expression levels of myosin IIIb and its ability to compensate for myosin IIIa need to be determined to elucidate why the recessive null mutations of myosin IIIa cause hearing loss only decades after birth and do not affect vestibular function.

References

Anderson DW, Probst FJ, Belyantseva IA, Fridell RA, Beyer L, Martin DM, Wu D, Kachar B, Friedman TB, Raphael Y, Camper SA (2000) The motor and tail regions of myosin XV are critical for normal structure and function of auditory and vestibular hair cells. Hum Mol Genet 9:1729–1738.

Belyantseva IA, Boker ET, Naz S, Frolokov Naz, Sellers JR, Ahmed ZM, Griffith AJ, Friedman TB (2005) Myosin-XVa is required for tip localization of whirlin and differential elongation of hair-cell stereocilia. Nat Cell Biol 7:148–156.

Berg JS, Cheney RE (2002) Myosin-X is an unconventional myosin that undergoes intrafilopodial motility. Nat Cell Biol 4:246–250.

Boeda B, El-Amraoui A, Bahliou A, Goodyear R, Caviet L, Blanchard S, Perfettini L, Fath KR, Shorte S, Reiners J, Houdusse A, Lepagin P, Wolfurm U, Richardson G, Petit C (2002) Myosin VIIa, harmonin and cadherin 23, three Usher I gene products that cooperate to shape the sensory hair cell bundle. EMBO J 21:6689–6699.

Cronin MA, Diao F, Tsunoa S (2004) Light-dependent subcellular translocation of Gqα-phlin in Drosophila photoreceptors is facilitated by the photoreceptor-specific myosin III NINAC. J Cell Sci 117:4797–4806.

Dose AC, Burnside B (2000) Cloning and chromosomal localization of a human class III myosin. Genomics 67:335–342.

Dose AC, Burnside B (2002) A class III myosin expressed in the retina is a potential candidate for Bardet-Biedl syndrome. Genomics 79:621–624.

Dose AC, Hillman DW, Wong C, Sohblng L, Lin-Jones J, Burnside B (2003) Myo3a, one of two class III myosins genes expressed in vertebrate retina, is localized to the caycal processes of rod and cone photoreceptors and is expressed in the sacculus. Mol Biol Cell 14:1058–1073.

Dumont RA, Zhao YD, Holt JR, Bahler M, Gillespie PG (2002) Myosin-I isoforms in neonatal rodent auditory and vestibular epithelia. J Assoc Res Otolaryngology 3:373–389.

Faix J, Rottner K (2006) The making of filopodia. Curr Opin Cell Biol 18:18–25.

Garcia JA, Yee AG, Gillespie PG, Corey DP (1998) Localization of myosin-Id near both ends of tip links in frog saccular hair cells. J Neurosci 18:8637–8647.

Gillespie PG, Cyr JL (2004) Myosin-1c, the hair cell’s adaptation motor. Annu Rev Physiol 66:521–545.

Gillespie PG, Dumont RA, Kachar B (2005) Have we found the tip link, transduction channel, and gating spring of the hair cell? Curr Opin Neurol 15:389–396.

Grati M, Schneider ME, Lipkow K, Strehler EE, Wenthold RJ, Kachar B (2006) Rapid turnover of stereocilia membrane proteins: evidence from the trafficking and mobility of plasma membrane Ca2+/ATPase 2. J Neurosci 26:6386–6395.

Kachar B, Parakkal M, Kurc M, Zhao Y, Gillespie PG (2000) High-resolution structure of hair-cell tip links. Proc Natl Acad Sci USA 97:13336–13341.

Kaltenbach JA, Falzarano PR, Simpson TH (1994) Postnatal development of the hamster cochlea. II. Growth and differentiation of stereocilia bundles. J Comp Neurol 350:187–198.

Komaba S, Inoue A, Maruta S, Hosoya H, Ikebe M (2003) Determination of human myosin II as a motor protein having a protein kinase activity. J Biol Chem 278:21352–21360.

Kros CJ, Marcotti W, van Netten SM, Seif van Netten, Libby RT, Brown SD,
Richardson GP, Steel KP (2002) Reduced climbing and increased slipping adaptation in cochlear hair cells of mice with Myo7a mutations. Nat Neurosci 5:41–47.

Lee SJ, Montell C (2004) Light-dependent translocation of visual arrestin regulated by the NINAC myosin III. Neuron 43:95–103.

Les Erickson F, Corsa AC, Dosé AC, Burnside B (2003) Localization of a class III myosin to filopodia tips in transfected HeLa cells requires an actin-binding site in its tail domain. Mol Biol Cell 14:4173–4180.

Li HS, Montell C (2000) TRP and the PDZ protein, INAD, form the core complex required for retention of the signalplex in Drosophila photoreceptor cells. J Cell Biol 150:1411–1422.

Liang Y, Wang A, Belyantseva IA, Anderson DW, Probst FJ, Barber TD, Miller W, Touchman JW, Jin L, Sullivan SL, Sellers JR, Camper SA, Lloyd RV, Kachar B, Friedman TB, Fridell RA (1999) Characterization of the human and mouse unconventional myosin XV genes responsible for hereditary deafness DFNB3 and shaker 2. Genomics 61:243–258.

Lin HW, Schneider ME, Kachar B (2005) When size matters: the dynamic regulation of stereocilia lengths. Curr Opin Cell Biol 17:55–61.

Lin-Jones J, Parker E, Wu M, Dosé A, Burnside B (2004) Myosin 3A transgene expression produces abnormal actin filament bundles in transgenic Xenopus laevis rod photoreceptors. J Cell Sci 117:5825–5834.

Ng KP, Kambara T, Matsuura M, Burke M, Ikebe M (1996) Identification of myosin III as a protein kinase. Biochemistry 35:9392–9399.

Philipp S, Flockerzi V (1997) Molecular characterization of a novel human PDZ domain protein with homology to INAD from Drosophila melanogaster. FEBS Lett 413:243–248.

Porter JA, Montell C (1993) Distinct roles of the Drosophila ninaC kinase and myosin domains revealed by systematic mutagenesis. J Cell Biol 122:601–612.

Ricci AJ, Kachar B, Gale J, Van Netten SM (2006) Mechano-electrical transduction: new insights into old ideas. J Membr Biol 209:71–88.

Rzadzinska AK, Schneider ME, Davies C, Riordan GP, Kachar B (2004) An actin molecular treadmill and myosins maintain stereocilia functional architecture and self-renewal. J Cell Biol 164:887–897.

Seiler C, Ben-David O, Sidi S, Hendrich Sidi, Rusch A, Burnside B, Avraham KB, Nicolson T (2004) Myosin VI is required for structural integrity of the apical surface of sensory hair cells in zebrafish. Dev Biol 272:328–338.

Sellers JR (2000) Myosins: a diverse superfamily. Biochim Biophys Acta 1496:3–22.

Sousa AD, Cheney RE (2005) Myosin-X: a molecular motor at the cell’s fingertips. Trends Cell Biol 15:533–539.

Stauffer EA, Scarborough JD, Hirono M, Miller H, Shah K, Mercer JA, Holt JR, Gillespie PG (2005) Fast adaptation in vestibular hair cells requires myosin-1c activity. Neuron 47:541–553.

Steyger PS, Gillespie PG, Baird RA (1998) Myosin Iβ is located at tip link anchors in vestibular hair bundles. J Neurosci 18:4603–4615.

Tilney LG, Tilney MS, Cotanche DA (1988) New observations on the stereocilia of hair cells of the chick cochlea. Hear Res 37:71–82.

Walsh T, Walsh V, Vreugde S, Hertzano R, Shahin H, Haika S, Lee MK, Kanaan M, King MC, Avraham KB (2002) From flies’ eyes to our ears: mutations in a human class III myosin cause progressive nonsyndromic hearing loss DFNB30. Proc Natl Acad Sci USA 99:7518–7523.

Wes PD, Xu XZ, Li HS, Chien F, Doberstein SK, Montell C (1999) Termination of phototransduction requires binding of the NINAC myosin III and the PDZ protein INAD. Nat Neurosci 2:447–453.

Wu DK, Oh SH (1996) Sensory organ generation in the chick inner ear. J Neurosci 16:6454–6462.

Zine A, Romand R (1996) Development of the auditory receptors of the rat: a SEM study. Brain Res 721:49–58.