Knockdown of ttc26 disrupts ciliogenesis of the photoreceptor cells and the pronephros in zebrafish

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ABSTRACT In our effort to understand genetic disorders of the photoreceptor cells of the retina, we have focused on intraflagellar transport in photoreceptor sensory cilia. From previous mouse proteomic data we identified a cilia protein Ttc26, orthologue of dyf-13 in Caenorhabditis elegans, as a target. We localized Ttc26 to the transition zone of photoreceptor and to the transition zone of cilia in cultured murine inner medullary collecting duct 3 (mIMCD3) renal cells. Knockdown of Ttc26 in mIMCD3 cells produced shortened and defective primary cilia, as revealed by immunofluorescence and scanning electron microscopy. To study Ttc26 function in sensory cilia in vivo, we utilized a zebrafish vertebrate model system. Morpholino knockdown of ttc26 in zebrafish embryos caused ciliary defects in the pronephric kidney at 27 h postfertilization and distension/dilation of pronephros at 5 d postfertilization (dpf). In the eyes, the outer segments of photoreceptor cells appeared shortened or absent, whereas cellular lamination appeared normal in retinas at 5 dpf. This suggests that loss of ttc26 function prevents normal ciliogenesis and differentiation in the photoreceptor cells, and that ttc26 is required for normal development and differentiation in retina and pronephros. Our studies support the importance of Ttc26 function in ciliogenesis and suggest that screening for TTC26 mutations in human ciliopathies is justified.

INTRODUCTION

Inherited retinal degenerations (IRDs) are important causes of blindness (Pierce, 2001). These disorders are characterized by dysfunc-

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Abbreviations used: BBS, Bardet-Biedl syndrome; bps, beats per second; BSA, bovine serum albumin; dpf, days postfertilization; EGFP, enhanced green fluorescent protein; PBS, phosphate-buffered saline; PSC, photoreceptor sensory cilium; RNA, RNA interference; RP, retinitis pigmentosa; RT-PCR, reverse transcriptase PCR; SEM, scanning electron microscopy; shRNA, short hairpin RNA; SSTR3, somatostatin receptor 3; TEM, transmission electron microscopy; TPR, tetratricopeptide repeats.

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phototransduction located in or associated with the membrane disks stacked in tight order at ~30 per micron along the axoneme. The transition zone in PCS is analogous to transition zones in other cilia and is where the triplet microtubule structure of the basal bodies converts to the doublet microtubule structure of the axoneme (Horst et al., 1990).

The recognition of photoreceptor outer segments as cilia connects retinal degenerative disorders such as retinitis pigmentosa (RP) and Leber congenital amaurosis (LCA) to other ciliary disorders. Mutations in genes that encode cilia proteins are increasingly recognized as common causes of disease. For example, inherited renal disorders, such as polycystic kidney disease and nephropathies, are now understood to be ciliopathies (Hildebrandt and Otto, 2005). Ciliary defects are also seen as the underlying cause of a number of inherited disorders that affect multiple organ systems. This helps explain the connection of retinal degeneration to ciliopathies, such as Bardet-Biedl syndrome (BBS), in which RP is found in association with multiple cilia-related disorders, including cystic renal disease, polydactyly, mental retardation, obesity and diabetes, gonadal malformations, and situs inversus (Satir and Christensen, 2007).

Ciliogenesis involves intraflagellar transport (IFT), which mediates the delivery of ciliary components to the growing ciliary tip, the point at which new cilia assembly occurs. The transported materials include tubulin subunits, tubulin-associated proteins, cell surface receptors, and other membrane proteins. IFT proteins have been biochemically characterized in Chlamydomonas as two complexes associated with anterograde (complex B) or retrograde (complex A) transport (Cole et al., 1998). Mutations in IFT complexes and other ciliary component genes have been identified in numerous ciliopathies (Pierce et al., 1999; Lehman et al., 2008; Louie et al., 2010; Muller et al., 2010; Davis et al., 2011).

The pleiotropic phenotypes of ciliary disorders suggest the involvement of genes having a highly conserved, essential function in the formation and/or maintenance of cilia. The inheritance patterns of ciliopathies are mostly autosomal recessive, but with extensive genetic heterogeneity (Badano et al., 2010; Davis et al., 2011). Mutations associated with anterograde (complex B) or retrograde (complex A) transport (Cole et al., 1998). Mutations in IFT complexes and other ciliary component genes have been identified in numerous ciliopathies (Pierce et al., 1999; Lehman et al., 2008; Louie et al., 2010; Muller et al., 2010; Davis et al., 2011).

Although some progress has been made in the characterization of IFT complexes and BBSome proteins, proteomic studies of several model systems indicate many proteins are expressed in cilia (Ostrowski et al., 2002; Keller et al., 2005; Pazour et al., 2005; Liu et al., 2007; Mayer et al., 2009; Jia et al., 2010). For example, we identified nearly 2000 proteins in PSC in proteomic analyses of mouse PSCs (Liu et al., 2007). These data suggest ciliogenesis is a complicated process and many uncharacterized proteins are involved.

In the work reported here, we have evaluated the role of a novel cilia protein Ttc26, which we detected in the mouse PSC proteome (Liu et al., 2007). Ttc26 has been classified as an IFT complex B protein based on in vitro coprecipitation with Ift88 (Follit et al., 2009). In addition, the Caenorhabditis elegans homologue of Ttc26, called daf-13, is proposed to interact with OSM-3, a homodimeric kinesin motor, to move cargo in the anterograde direction during IFT (Stanch et al., 1995; Blacque et al., 2005). In its N-terminal portion, Ttc26 contains several tetratricopeptide repeats (TPR). The TPR motif is enriched in the ciliary proteome and may mediate its interaction with other ciliary proteins in higher-order complexes. Notably, mutations in proteins with multiple TPRs, such as fleer/DYF-1, BBS4, BBS8, and TTC21B have been reported as ciliopathy-causing genes either in animal models or in human patients (Katsanis et al., 2002; Pathak et al., 2007; Harville et al., 2010; Riazuddin et al., 2010; Davis et al., 2011). The studies reported here demonstrate that Ttc26 is located in the transition zone of primary and photoreceptor cilia and is required for normal ciliogenesis in cultured cells and in vivo in zebrafish.

RESULTS
Ttc26 is a highly conserved protein expressed in tissues containing ciliated cells
We detected Ttc26 as a novel PSC protein in our proteomic analyses of PSCs isolated from mouse retina. In those analyses, Ttc26 was detected by eight unique peptides, suggesting that it is present in moderate abundance in mouse PSCs (Liu et al., 2007). To get an indication of the evolutionary importance of Ttc26, we searched the National Center for Biotechnology Information protein sequence database and found homologues from green algae, insects, and mammals. Alignment of Ttc26 homologue protein sequences from species related to the current study (CLUSTAL W method; Supplemen
tal Figure S1) revealed very high sequence conservation (humans and rodents: 97% identity, 99% similarity; zebrafish and humans/rodents: 80% identity, 90% similarity). Ttc26 is highly conserved across the species we examined. The sequence conservation is observed throughout the protein, not just in TPR domains.

We used northern blots to determine which mouse tissues express transcripts of Ttc26 and how abundant the transcripts are. In tests, which displayed the highest expression level, two different-sized transcripts were identified (Figure 1A). The larger transcript (4.2 kb) is consistent with the largest splice variant annotated in Ensembl (ENSMUST00000162554; CCDS51750) and was also detected in mouse retina, kidney, lung, and brain tissue. The expression level was low in liver and spleen, consistent with the low proportion of ciliated cells in these tissues (Figure 1A). The smaller transcript (2.5 kb) may represent an alternative splice variant in tests and is consistent with Ensembl transcript ENSMUST00000039394, in which exons 7–17 of the larger transcript are skipped. We also examined ttc26 developmental expression in embryonic zebrafish and found that ttc26 transcripts can be detected as early as the one-cell stage (possibly reflecting both maternal and embryonic contributions) and maintain a relatively stable level during the examined developmental stages (Figure 1B).

Ttc26 is localized in the transition zone of primary and photoreceptor sensory cilia
After confirming the temporal and spatial sequence of Ttc26 expression, we examined its subcellular localization by expressing a V5-tagged mouse Ttc26 fusion protein in murine inner medullary collecting duct 3 cells–somatostatin receptor 3–enhanced green fluorescent protein (mIMCD3-SSTR3-EGFP) cells. This stable cell line, derived from medullary collecting duct cells, expresses a green fluorescent protein (GFP) fused to SSTR3, a ciliary membrane receptor. By performing colocalization of Ttc26 with this ciliary marker, we showed that Ttc26 was concentrated in the ciliary base with gradually decreasing staining intensity toward the ciliary tip in interphase cells (Figure 1C). By immunostaining with antibodies to CEP164, a basal body distal appendage marker protein (Graser et al., 2007), we refined the Ttc26 subcellular location at the base to the ciliary transition zone. Interestingly, Ttc26 does not localize to the distal appendage itself, but rather lies immediately adjacent to it. However, the punctate signal spreading in whole cilia (Figure 1C) suggests that Ttc26 is not restricted to the transition zone and is possibly mobile in a protein complex.

We also studied Ttc26 subcellular location in neonatal rat photoreceptor cells using an in vivo retinal electroporation approach to
transfect the same plasmid DNA used in the mlMCD3 cell transfection. Using immunostaining to the V5 tag on the Ttc26 fusion protein, we were able to localize Ttc26 to the transition zone in photoreceptor cells (Figure 1D) and thus confirm it as a bona fide ciliary protein.

RNA interference (RNAi) knockdown of Ttc26 causes a defect in primary cilia in cultured kidney epithelial cells

The identification of Ttc26 as a transition zone protein suggests that it may play a functional role in ciliogenesis. To test this hypothesis, we used a ciliated renal epithelial cell model (mlMCD3) in vitro to study the effects of knockdown of Ttc26 expression. Three short hairpin RNA (shRNA) constructs directed at Ttc26 were introduced into target cells using a vector expressing GFP (pCAG-miR30-IRES-EGFP). This allowed us to identify all shRNA-transfected cells.

Cotransfection of these Ttc26 shRNAs with the V5-Ttc26 cDNA in CHO cells demonstrated that all three shRNA constructs had knockdown efficiencies of over 90% compared with control shRNAs (Figure 2A). Next we tested the effect of shRNA-mediated Ttc26 knockdown on ciliogenesis in mlMCD3 cells. In this case, cilia were detected by immunostaining with antibody to acetylated α-tubulin, an axoneme marker. Transfected mlMCD3 cells (green) were

FIGURE 1: Expression of Ttc26 in cultured mlMCD3 cells, rodent tissues, and zebrafish embryos. (A) Northern blot of Ttc26 in adult mouse tissues. Ttc26 expression is high in the testis, with a 2.5-kb transcript exclusively seen in testis and a 4.2-kb transcript detected in testis, brain, heart, lung, kidney, and retina. The 11.4-kb band may represent unprocessed RNA. (B) Developmental expression of ttc26 in zebrafish embryos. RT-PCR data revealed ttc26 transcription was active at the one-cell stage and maintained at a relatively stable level through the developmental process (note that samples of one-cell and four-cell embryos were not loaded on the same gel). (C) Top, V5-Ttc26 fusion protein (red) colocalized with cilia (green) in SSTR3-EGFP-mlMCD3 cells. Ttc26 protein is most concentrated in the ciliary base. Middle, Cep164 (red), a distal appendage protein in basal body, is expressed in cilia (green) of SSTR3-EGFP-mlMCD3 cells. Bottom, V5-Ttc26 protein is partially overlapped with Cep164 in the basal bodies of mlMCD3 cells. Right, merged images. Note that the two proteins locate closely but not overlap. Insets, enlarged images of the cilia.

(D) Neonatal rat photoreceptor cells transfected by in vivo electroporation with the plasmid pCAG-V5-Ttc26-IRES-EGFP are revealed by IRES-EGFP expression (green). Ttc26 recombinant protein, detected by immunostaining of the V5 tag, is localized to the transition zone of the transfected photoreceptor cells (red, arrows) on the top of the inner segments filled with water-soluble EGFP. Right, merged images.
exons 1 and 2, leading to the inclusion of intron 1 in the altered mRNA transcript. This introduces a premature termination codon 20 nucleotides into intron 1, which leads to a truncated protein and loss of function. Molecular analysis of reverse transcription PCR (RT-PCR) products from embryos injected with MO-SP revealed missplicing events (Figure 3, D and E) consistent with a loss-of-function phenotype similar to that caused by MO-AUG. Embryos injected with a control morpholino were normal in appearance (Figure 3A, 1 and 2). However, the morphant larvae exhibited body axis curvature (mostly ventral and sometimes dorsal) and kinked tails (Figure 3A, 3–10). The MO-AUG and MO-SP morpholinos led to very similar phenotypes in appearance and in histology, except the MO-SP morphants also displayed severe body edema. The histology of 5 d postfertilization (dpf) zebrafish eye morphogenesis in control and ttc26 morphants is presented in Figure 3B. At higher magnification (60×), the five principal laminae—three cellular and two plexiform layers—can be seen to be fully formed in the retinas of both control and morphant larvae, and the proliferative marginal zones (MZ) developed normally and remained nonlaminated in both controls and morphants. A higher-magnification view of the control retina shows clear morphological differentiation of photoreceptors with well-defined inner and outer segments at 5 dpf (Figure 3B). In contrast, both types of morphants revealed a defective photoreceptor ciliogenesis with only a few outer segments visible. Outer segments in these retinas were smaller, shortened, and disorganized. observed to have shortened cilia or missing cilia (arrows in Figure 2B). We further analyzed the images by quantifying the number of transfected cells with and without cilia and by measuring the ciliary length in both transfected cells and adjacent nontransfected cells. Using this approach, we found that knockdown of Ttc26 was associated with significantly reduced ciliary length (p < 0.01; Figure 2, B and C). Nontransfected mIMCD3 cells are shown for comparison in Figure 2D. To investigate the effects of Ttc26 knockdown on ciliary morphology, we examined shRNA-transfected and nontransfected mIMCD3 cells by scanning electron microscopy (SEM) and observed that cells with Ttc26 knockdown displayed shortened cilia with enlarged ends (Figure 2E).

**Morpholino knockdown of ttc26 disrupts photoreceptor outer segment morphogenesis in developing retina**

To further assess the functional importance of Ttc26, we turned to a vertebrate model organism, the zebrafish. Over a period of 2 d, the optic cup transforms from a single neuroepithelial sheet into a functional multilayered retina (Fadool and Dowling, 2008). This makes it particularly suitable for a developmental study of photoreceptor cell morphogenesis. We used knockdown by two ttc26 morpholino oligonucleotides (MO-AUG and MO-SP) to further test the hypothesis that ttc26 functions in ciliogenesis. The MO-AUG and MO-SP morpholinos were designed to interfere with translation initiation and RNA splicing, respectively. The MO-SP blocks splicing between exons 1 and 2, leading to the inclusion of intron 1 in the altered mRNA transcript. This introduces a premature termination codon 20 nucleotides into intron 1, which leads to a truncated protein and loss of function. Molecular analysis of reverse transcription PCR (RT-PCR) products from embryos injected with MO-SP revealed missplicing events (Figure 3, D and E) consistent with a loss-of-function phenotype similar to that caused by MO-AUG.

Embryos injected with a control morpholino were normal in appearance (Figure 3A, 1 and 2). However, the morphant larvae exhibited body axis curvature (mostly ventral and sometimes dorsal) and kinked tails (Figure 3A, 3–10). The MO-AUG and MO-SP morpholinos led to very similar phenotypes in appearance and in histology, except the MO-SP morphants also displayed severe body edema. The histology of 5 dpf photoreceptor eye morphogenesis in control and ttc26 morphants is presented in Figure 3B. At higher magnification (60×), the five principal laminae—three cellular and two plexiform layers—can be seen to be fully formed in the retinas of both control and morphant larvae, and the proliferative marginal zones (MZ) developed normally and remained nonlaminated in both controls and morphants. A higher-magnification view of the control retina shows clear morphological differentiation of photoreceptors with well-defined inner and outer segments at 5 dpf (Figure 3B). In contrast, both types of morphants revealed a defective photoreceptor ciliogenesis with only a few outer segments visible. Outer segments in these retinas were smaller, shortened, and disorganized.
Transmission electron microscopy (TEM) images of the controls show well-formed photoreceptor outer segments. In contrast, morphant eyes reveal missing and ill-formed outer segments (Figure 3C).

**MO-knockdown of ttc26 results in dilated pronephric tube and duct**

The observation of edema in ttc26 morphants prompted us to look at the pronephros, the embryonic kidney, of zebrafish larvae for possible defects in fluid flow and in biogenesis of cilia, including motile cilia. Indeed, as shown in Figure 4, we found that morpholino knockdown of ttc26 was associated with defects in cilia in the pronephros as early as 27 h postfertilization (hpf), with disrupted and disorganized cilia (Figure 4A). Video images taken at 2 dpf clearly show enlarged pronephric ducts and disruption of multiciliated-cell cilia beat coordination (Supplemental Movies S1, S2, and S3). The recorded video data showed the multiciliated-cell cilia beat coordination is disrupted. The overall beat rates increase from 34 beats per second (bps; control) to ∼60 bps (morphants), most likely due to tubule lumen dilation (Hellman et al., 2010; Figure 5). The distended and dilated pronephric tubules/ducts were observed in many morphants that exhibited precardiac and body edema at 5 dpf (Figure 4, C–E).

The finding that both the MO-AUG and MO-SP morpholinos produced the same phenotypes in both eyes and pronephric kidney supports the conclusion that these results are due to genuine ttc26 knockdown and not to an off-target effect. To confirm the predicted phenotypes of ttc26 morphant larvae, we examined eye and pronephric kidney morphology. Zebrafish embryos were injected with two ttc26 MOs (MO-AUG and MO-SP) or control MO. (A) Knockdown of ttc26 results in developmental defects. Panel 1, lateral view of 120-hpf larva injected with control MO. Panel 2, dorsal view of the control MO-treated larva. Panels 3–6, lateral views of 120-hpf morphant larvae injected with ttc26 morpholino MO-AUG. The ttc26 morphants displayed curled or kinked tails, precardiac edema, and shortening of the body. Panels 7–10, dorsal or lateral view of 72-hpf morphant larvae injected with ttc26 morpholino MO-SP. The curled or kinked tails and body curvature are very similar to those of MO-AUG morphants. (B) Eye morphology is altered in ttc26 morphants at 5 dpf. Histological sections are shown for eyes, central retina (CR), and retinal marginal zone (MZ) from larvae injected with control MO and ttc26 MO-AUG. Left, retina of 5-dpf fish injected with control MO display well-formed laminae (top), with details visible in high magnification (60×) views of the central retina (CR, middle) and marginal zone (MZ, bottom). The higher magnification (60×) views of the morphant retinas reveal shortened and disorganized photoreceptor outer segments in the retina, with otherwise normal lamination. (C) TEM of the control-MO larvae at 5 dpf shows normally developed photoreceptor outer segments with densely packed disks (left). TEM of the morphant larvae shows that outer segments are smaller, shortened, and disoriented (right), consistent with the morphology seen in the light micrographs in (B). Note that a disorganized partial outer segment can be seen on the left adjacent to a cone photoreceptor (red arrow). N, nucleus; M, mitochondria; OS, outer segment. (D) RT-PCR analyses confirm that morpholino MO-SP interferes with splicing of ttc26 mRNA in larvae. Forty-eight hours postfertilization and 120-hpf larvae injected with control-MO or MO-SP were used for RNA extraction and RT-PCR. The lower band (590 base pairs) visible in both conditions represents PCR product from correctly spliced mRNA. The larger PCR product (1553 base pairs, containing the 963–base pair nonsplicing intron 1) is detected after MO-SP treatment due to the blocked splice donor site. As shown, although not complete, splice blocking is more evident at 48 hpf, consistent with some recovery from the morpholino effect at 120 hpf. (E) The binding positions of the PCR primers (above) and the MO-SP oligo (below) are illustrated in a diagram. For simplicity, only exons 1 and 2 are shown.
The data presented here indicate that Ttc26 is localized to cilia in both mouse and rat cells and is developmentally required for ciliogenesis in zebrafish photoreceptor cells and pronephric kidney. Ttc26 is localized in the primary cilia and enriched in the transition zone of the mIMCD3 and rodent photoreceptor cells. Knockdown of Ttc26 in mIMCD3 cells produced shortened and defective cilia. Furthermore, morpholino knockdown of ttc26 in zebrafish caused larvae to exhibit absent or deformed photoreceptor outer segments, but with normal lamination in the morphant retina. The morphant larvae also developed defective motile cilia in the pronephros. These morphant phenotypes demonstrate that ttc26 is required for ciliogenesis and normal ciliary function.

FIGURE 4: Morpholino knockdown of ttc26 disrupts motile cilia and causes tubule dilation in zebrafish pronephros. (A) Cilia in the anterior and posterior regions of the pronephric kidneys in larvae at 27 hpf. Green, cilia revealed by immunostaining of acetylated α-tubulin; blue, nuclei. Top, cilia are long and well-organized in larvae injected with the control MO oligo. Middle and bottom, knockdown of ttc26 by transfection with MO-AUG or MO-SP morpholinos leads to shortened, disorganized, and disoriented cilia. (B) Kidney cross-section in a 5-dpf larva shows normal morphology after injection with control-MO (C–E) Injection with MO-AUG or MO-SP morpholino leads to distended/dilated pronephric tubes and ducts. The zebrafish larvae used are shown below the corresponding micrograph images, with the defects marked (*). gl, glomerulus; pt, pronephric tubule; pd, pronephric duct.

that the morphological effects in zebrafish larvae were indeed associated with knockdown of ttc26, we used RT-PCR analysis to analyze the mRNA present in control and morphant larvae. Morpholino MO-SP treatment of zebrafish larvae partially blocks splicing of ttc26 mRNA, and thus leads to a larger-sized PCR product (Figure 3, D and E). To further demonstrate the specificity of the MO-induced knockdown phenotypes, we performed rescue experiments with coinjection of ttc26-MO and Ttc26 mRNA. Coinjection of 66 ng of mouse Ttc26 mRNA with MO-AUG reduced the percentage of abnormal larvae from 95% (21/22) in the embryos injected with MO-AUG alone to 51% (23/45) in the cojected embryos (p < 0.001).

DISCUSSION
The data presented here indicate that Ttc26 is localized to cilia in both mouse and rat cells and is developmentally required for kidney cysts, randomized left–right asymmetry, hydrocephalus, and rod outer segment defects in zebrafish (Pathak et al., 2007).

Related genes involved in ciliary defects
The highly conserved polypeptide sequence of Ttc26, both within its TPR domains and in other regions, suggests an important function for this protein. Furthermore, ttc26 knockdown in the eye and pronephros led to phenotypes that match those observed with defects in other ciliary genes in zebrafish larvae. These include body curvature, cardial edema, pronephric distention and dilation, and photoreceptor outer segment defects (Malicki et al., 2011). The overlap in phenotypes implies that ttc26 might function similarly to other TPR domain proteins found in ciliary protein complexes.

Ciliopathy-causal mutations have been identified in several TPR domain-containing genes, including BBS4, BBS8, TTC21B, and TTC30B (Katsanis et al., 2002; Billingsley et al., 2010; May-Simera et al., 2010; Razuddin et al., 2010; Davis et al., 2011). BBS4 and BBS8 are key scaffold components of the BBSome ciliary protein transport complex. TTC21B has been localized to centrosomes and is required for retrograde IFT, and causal mutations have been identified in human patients (Davis et al., 2011). Fleer/dfy-1/TTC30B is a regulator of ciliary tubulin polyglutamylation. Mutants in *fleer* showed
Zebrafish morphants lacking ttc26 are viable at least to the 5-dpf stage and show photoreceptor cells with short or missing PSCs. The loss of ttc26 appears to prevent or delay PSC formation during early photoreceptor cell differentiation. While outer segment disk material is generated in some photoreceptor cells, it does not appear that this can be organized into formed PSCs in most photoreceptor cells. Although the morphant photoreceptor cells displayed defects in PSC formation, they maintained the ability to laminate and differentiate normally in the retina.

Relation of kidney defects to ciliopathy and ttc26 function
A hallmark of many human ciliopathies is the formation of kidney cysts, which often begins during fetal development. The observation of edema in zebrafish ttc26 morphants prompted us to look for defects in cilia early in pronephros at 27 hpf. The kidney is a vital organ controlling homeostasis of water and electrolytes, osmoregulation of body fluids, and filtration of toxic metabolic waste products. Cilia in zebrafish pronephros have been reported to be motile and to drive fluid flow (Kramer-Zucker et al., 2005). Therefore disruption of the cilia would logically be accompanied by a loss of fluid flow, leading to fluid accumulation and subsequent pronephric distension and dilation. As expected, ttc26 morphant larvae at 27 hpf exhibited disorganized and disrupted cilia, loss of cilia bundles in the pronephric duct, and distended pronephric tubules and ducts, and this would logically contribute to the disruption of kidney function and severe edema observed in zebrafish larvae after ttc26 knockdown.

In summary, our characterization of Ttc26 localization and function in mouse and rat tissue, mIMCD3 cells, and in zebrafish provides the first comprehensive studies of Ttc26 gene expression and function in primary and sensory cilia. This work demonstrates that Ttc26 plays a critical role in ciliogenesis and suggests that loss of Ttc26 function could be implicated in retinal, renal, and other disease states in humans. Further studies confirming and extending these findings in other species will help us to better understand the function of Ttc26 in cilia. In this light, it seems logical to screen human patient populations with ciliopathies for mutations of TTC26 and genes that encode other ciliary proteins involved in IFT.

MATERIALS AND METHODS
Gene expression reagents
Mouse Ttc26 cDNA was amplified by RT-PCR from mouse retina, cloned into a pENTR/D-TOPO entry vector (Invitrogen, Carlsbad, CA), and fully sequence-verified. The coding sequence was moved by recombination to a Gateway destination expression vector modified to contain an N-terminal V5 epitope tag in-frame (pCAG-V5-IRES-EGFP; Hartley et al., 2000; Matsuda and Cepko, 2004, 2007). The sequences of shRNA knockdown reagents were generated using the RNAi Central website (Cold Spring Harbor, http://hannonlab.cshl.edu). Oligonucleotides of shRNA (obtained from IDT, Coralville, IA) were PCR-amplified and cloned into a pCAG-miR30-IRES-EGFP vector, modified by our laboratory for this purpose using established methods (Hartong et al., 2006). Plasmid DNA was purified using the EndoFree Plasmid Maxi kit (Qiagen, Valencia, CA).

Cell culture and transfection
An mIMCD3 stable cell line expressing SSTR3-EGFP, a ciliary membrane marker (gift of Gregory J. Pazour, University of Massachusetts Medical School), was used for Ttc26 subcellular localization. CHO-K1 and wild-type mIMCD3 cell lines were purchased from the American Type Culture Collection. The mIMCD3 cells were maintained in DMEM:F12 media supplemented with 10% fetal bovine serum (FBS) and 0.5 mM sodium pyruvate (Invitrogen). CHO cell culture was performed in F-12K medium (Invitrogen) supplemented with 10% FBS. Transfection was performed with Lipofectamine 2000 or Lipofectamine LTX reagent (Invitrogen) using 4 μg DNA per well and cells at 60–70% confluency in a six-well plate with glass coverslips (25CIR-1D; Fisher, Pittsburgh, PA). The cells were processed for immunocytochemistry at 72 h after transfection. Detection of the recombinant protein fused to the V5 epitope tag was accomplished by immunocytochemical detection or immunoblot of the V5 tag.

Antibody reagents
Monoclonal antibody (mAb) against acetylated α-tubulin (clone 6-11B-1) was obtained from Sigma-Aldrich (St. Louis, MO) and anti-V5 tag mAb from Invitrogen. Anti–CEP164 antibody (human), recognizing a basal body distal appendage marker protein, was a gift (Graser et al., 2007). Anti-β-actin antibody (C-11) was purchased from Santa Cruz Biotechnology (Santa Cruz, CA). All secondary antibodies conjugated with fluorophores were from Invitrogen, except the antibody used for immunoblot, which was obtained using the EndoFree Plasmid Maxi kit (Qiagen, Valencia, CA).
from LI-COR (Lincoln, NE). The chemical reagents were purchased from Sigma-Aldrich.

**Immunofluorescence microscopy**

All incubations and washes were carried out in 1× phosphate-buffered saline (PBS) at room temperature. Cells were fixed in 4% paraformaldehyde for 10 min, permeabilized with 0.5% Triton X-100 in PBS for 10 min, and blocked with 1% bovine serum albumin (BSA) and 0.2% Triton X-100 in PBS for 10 min. Next the cells were incubated with antibody to the V5 expression tag (1:2500 dilution) in blocking buffer for 1 h at room temperature. After three 10-min washes in PBS, a secondary antibody incubation was carried out for 1 h in Alexa Fluor 555–conjugated goat anti–mouse immunoglobulin G (IgG; 1:1000 dilution). The cells were then washed once in PBS for 10 min, incubated with Hoechst dye (Invitrogen) at 1:1000 dilution for 5 min, and washed twice (10 min each wash). Forocolocalization to the basal body, cells were incubated using the above procedure with antibody against human CEP164 for 1 h (1:1000 dilution) and Alexa Fluor–conjugated goat anti–mouse IgG (1:1000 dilution; Invitrogen) for 1 h. The coverslips were mounted using Fluoromount (Electron Microscopy Sciences, Hatfield, PA). Fluorescence signals were visualized using a Nikon TE300 fluorescence microscope.

**In vivo electroporation**

To localize Ttc26 expression in photoreceptor cells, we used an in vivo electroporation technique in rat photoreceptor cells. Briefly, 0.5 μl of endotoxin-free pCAG-V5-Ttc26-IREGFP plasmid was injected subretinally into the right eyes of neonatal rats. Injected plasmid was electroporated into retinal cells using tweezer-type electrodes as previously described (Matsuda and Cepko, 2004). The plasmid was injected into four to six eyes to provide sufficient retina with both good transfection and morphology for analysis. Animals were killed 4 wk following injection, and frozen sections were prepared from the portions of the eyecups with EGFP signal. Sections were immunostained as described previously using anti-V5 antibody followed by Alexa Fluor 555–conjugated secondary antibody. The location of V5-tagged proteins in photoreceptor cells was evaluated by three-dimensional reconstructions of the confocal image stacks generated and analyzed using Velocity 3D imaging software (Mountain View, CA).

**Northern blot and immunoblot analyses**

RNA expression levels were analyzed by northern blot analysis. To prepare a cDNA probe for northern blot analysis, we cloned the full-length mouse Ttc26 cDNA (NM_153600) into a pCR2-TOPO vector. The plasmid was digested with EcoRI, and the insert DNA was purified using the QIAEX II gel extraction kit (Qiagen). Northern blot analysis was performed as described previously (Zhang et al., 2000). Briefly, total RNA (25 μg) was extracted with TRIzol reagent (Invitrogen) and transferred to a Gene Screen Plus nylon membrane (Perkin Elmer–Nycomed, New York) and hybridized with the RNA blot at 10°C in hybridization buffer. The blots contained multiple tissues, including retina from a wild-type adult mouse. For this reason, after decay of the cDNA probe signals, the membranes were rehybridized with an 32P-labeled cDNA probe for 18S RNA to assess and normalize RNA sample loading. The signal was scanned using a Typhoon 9400 phosphorimager (GE, Waukesha, WI).

Immunoblot analysis was used to verify knockdown of gene constructs expressed in CHO cells. Cells were transfected with pCAG V5-Ttc26-IREGFP plasmid DNA (4 μg well) in a six-well plate. At 48 h after transfection, the plate was rinsed twice with cold 1× PBS and then the cells were lysed immediately with 2x LDS sample buffer (Invitrogen) and sonicated briefly on ice. Total protein (100 μg/well) was separated on a precast NuPAGE 4–12% Bis-Tris Gel (Invitrogen) and transferred to a low-fluorescence polyvinylidene fluoride membrane (GE, Waukesha, WI). The membrane was incubated with Odyssey blocking buffer (LI-COR, Lincoln, NE) for 1 h at room temperature. The membrane was incubated with monoclonal anti-V5 antibody (1:5000 dilution) overnight with gentle rocking in a cold room, washed 4 times with 1× PBS with 0.1% Tween-20, incubated with IRDye (LI-COR) goat anti-mouse IgG (1:10,000 dilution) for 1 h at room temperature, and washed three times with PBS with 0.1% Tween-20. After a final wash with 1× PBS, the signals were detected with an Odyssey infrared imager (LI-COR). The loading control antibody to β-actin was used at 1:1000. Protein levels (expressed in fluorescence units) were quantified with software provided by LI-COR.

**MOs and zebrafish embryo injection**

Wild-type zebrafish (Danio rerio) of the AB strain were obtained from the University of Pennsylvania zebrafish core facility (Philadelphia, PA). Embryos were collected in E3 solution after natural spawns and kept at 28.5°C in an incubator using standard protocols (Westerfield, 2007). Morpholinos designed to block ttc26 RNA translation (MO-AUG, 5′-TCGCGCTTCATCCGAGACAGAGCAT-3′) or to block splicing (MO-SP, 5′-ATATGTTGCTTTGATGCACCT-GTT-3′) and a standard control oligo (5′-CTCTTACCTCAGTTACAATTATA-3′) were obtained from Gene Tools (Philomath, OR).

Embryos were injected at the one- to two-cell stages with 0.35–0.45 mM MO (3.5–4.5 ng of ttc26 MO or 4.5 ng of control MO). Embryos were imaged with a Stemi SV11 stereomicroscope (Zeiss) and photographed with an AxioCam digital camera (Zeiss), and the images were analyzed with AxioVision software (Zeiss). Embryos were prepared for flat-mounting by dissecting the yolk away from the embryo proper with a pair of fine forceps; the embryos were then mounted between two glass coverslips. Immunofluorescence detection of acetylated tubulin in the whole-mount embryos was performed according to a published protocol (Jaffe et al., 2010).

High-speed video microscopy analysis has been described in detail in a recent publication (Pathak et al., 2011). Larvae at 2 dpf were used for the video recording.

**RT-PCR**

Whole embryos (both morphant and control) collected at different developmental stages were used for total RNA extraction with TRIzol reagent (Invitrogen). Total RNA (2 μg per sample) was reverse-transcribed using the Superscript first-strand synthesis system for RT-PCR (Invitrogen). The following primer pairs were used for RT-PCR. For ttc26 developmental expression: forward, 5′-AT-GCGCTTCTATACAGGAAGGC-3′; reverse, 5′-AGAGGCGCATG-CCTGTAGCCGGCT-3′. For the phenotype rescue studies, mouse Ttc26 cDNA was amplified by PCR with a pair of primers (forward, 5′-TGCGGCGGATCCTGAGGGCG-3′; reverse, 5′-ATGACAACTAGGACGCGAAGCG-3′) and cloned into an Sp6/T7 promoter vector (pCS2P+; Addgene, Cambridge, MA). The plasmid was linearized with SnaBl. Capped Ttc26 mRNA was synthesized in vitro using the...
mMESSAGE mMACHINE SP6 kit (Ambion, Austin, TX) and purified by the RNeasy Plus micro kit (Qiagen). Mouse Ttc26 mRNA (5–17 pg) was co-injected with MO-AUG (0.35 ng) into one- to two-cell stage zebrafish embryos.

Histology and SEM and TEM
Zebrafish embryos were processed for histology and TEM following a Cold Spring Harbor Protocol (http://cshprotocols.cshlp.org/content/2007/6/pdb.prot4772.abstract). For general histology, transverse semithin sections (1 μm) were stained with alkaline Toluidine blue. For TEM, thin sections (90 nm) were mounted on electron microscopy grids and stained with lead citrate and uranyl acetate. TEM images were collected on a FEI-Tecnai T12 transmission electron microscope. For SEM, cultured cells were processed using standard protocols. Briefly, mIMCD3 cells grown on 13-mm coverslips were rinsed twice with PBS, washed three times with 50 mM Na-cacodylate buffer (pH 7.3), fixed for 2 h with 2% glutaraldehyde in 50 mM Na-cacodylate buffer, and dehydrated in a graded series of ethanol concentrations over a period of 1.5 h. Dehydration in 100% ethanol was done three times. The dehydrated samples were immersed twice for 10 min in 100% hexamethyldisilazane (HMDS; Sigma-Aldrich, St. Louis, MO), which was followed by air-drying for 30 min (Braet et al., 1997). The samples were mounted on stubs and sputter-coated with gold palladium. Specimens were observed and photographed using a Philips XL20 scanning electron microscope (FEI, Hillsboro, OR) at 10-kV accelerating voltage.

Statistical analyses
Statistical analyses were performed with Minitab software. A Student’s t test with unequal variance or chi-square test was used for the analyses.

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