The ciliary transition zone functions in cell adhesion but is dispensable for axoneme assembly in C. elegans

Clementine Schouteden,* Daniel Serwas,* Mate Palfy, and Alexander Dammermann

Max F. Perutz Laboratories, University of Vienna, Vienna Biocenter (VBC), A-1030 Vienna, Austria

Cilia are cellular projections that perform sensory and motile functions. A key ciliary subdomain is the transition zone, which lies between basal body and axoneme. Previous work in Caenorhabditis elegans identified two ciliopathy-associated protein complexes or modules that direct assembly of transition zone Y-links. Here, we identify C. elegans CEP290 as a component of a third module required to form an inner scaffolding structure called the central cylinder. Co-inhibition of all three modules completely disrupted transition zone structure. Surprisingly, axoneme assembly was only mildly perturbed. However, dendrite extension by retrograde migration was strongly impaired, revealing an unexpected role for the transition zone in cell adhesion.

Introduction

The cilium is an evolutionarily ancient structure found in most eukaryotic lineages (Hodges et al., 2010). Projecting from the cell surface, cilia mediate sensory perception and development signaling. A subset of cilia is also motile, enabling movement of cells and fluids. Cilia dysfunction has been linked to a spectrum of human disorders, known as ciliopathies (Fliegauf et al., 2007), ranging from disorders affecting a single type of cilium to broad-based disorders affecting multiple tissues and organs.

All cilia arise from a centriole-derived basal body, connected to the plasma membrane via transition fibers. The axonemal microtubules constituting the core of the cilium are extensions of basal body microtubules. Immediately distal to the basal body lies the transition zone, distinguished by several conserved features, including a central cylinder or apical ring that lies internal to the microtubule doublets and Y-links connecting axonemal microtubules to the ciliary membrane (Reiter et al., 2012). The ciliary membrane is contiguous with the plasma membrane but is compositionally distinct, as is the underlying cytoplasm, suggesting that structures at the ciliary base (the transition fibers and/or transition zone) control access to the ciliary compartment (Rosenbaum and Witman, 2002).

During ciliogenesis, centrioles dock to a membrane, either to a vesicle in the cytoplasm that fuses with the plasma membrane (Sorokin, 1962) or to the plasma membrane itself. Docking requires the transition fibers (Schmidt et al., 2012), which in vertebrates are derived from appendages present on mature centrioles. Finally, motor-driven intraflagellar transport (IFT; Cole et al., 1998) extends the ciliary axoneme. While the molecular mechanisms underlying axoneme extension are comparatively well understood, less is known about basal ciliary structures and their role in the early stages of ciliogenesis (Reiter et al., 2012). For the transition zone, proteomic approaches in vertebrates identified three distinct ciliopathy-associated multiprotein complexes or modules: the MKS, NPHP-1,-4,-8 (NPHP) and NPHP-5,-6 (CEP290) modules (Garcia-Gonzalo et al., 2011; Sang et al., 2011; Chih et al., 2012). These are supported by genetic analyses in Caenorhabditis elegans, which have established independent assembly pathways for two modules (MKS and NPHP; Williams et al., 2011). It is currently unclear where these modules fit within the transition zone structure or how it assembles during ciliogenesis. Further, while the transition zone clearly functions as a ciliary gate, there is some uncertainty as to what extent it is also required for axoneme assembly, with loss of transition zone components in vertebrates resulting in loss of cilia in some tissues but not others (Garcia-Gonzalo et al., 2011; Sang et al., 2011; Chih et al., 2012).

The nematode C. elegans has emerged as a major experimental model to study centrioles and cilia. A key feature of C. elegans is that cilia are limited to the dendritic endings of postmitotic sensory neurons and are dispensable for viability and fertility, which facilitates loss-of-function studies (Inglis et al., 2007). Here, we take advantage of this experimental model to dissect the assembly and function of the transition zone. We identify a C. elegans homologue of CEP290 and show it to be a component of a third, independently targeted transition zone module required to form the central cylinder, which acts as an inner scaffolding structure for transition zone assembly in C. elegans.
Figure 1. **CCEP-290 belongs to a third transition zone module required for assembly of the central cylinder.** (A) Transition zone modules in *C. elegans*, based on genetic interactions and localization interdependencies (Huang et al., 2011; Williams et al., 2011; this study). (B) Modules in vertebrates, based on proteomic studies (green, red, blue: Sang et al., 2011; dark green: Chih et al., 2012; light green: Garcia-Gonzalo et al., 2011). *, *C. elegans* names.
assembly. Simultaneous inhibition of all three modules completely disrupts transition zone structure. Surprisingly, axoneme assembly is not significantly impaired. Instead, perturbing the transition zone disrupts cell–matrix interactions during dendrite extension, revealing an unexpected role for the transition zone in cell adhesion.

Results and discussion
Increasing evidence points to the transition zone being organized into multiple protein complexes or modules with distinct functions. Best characterized are the MKS and NPHP modules, which are composed of proteins mutated in the ciliopathies Meckel syndrome and nephronophthisis. Previous work in C. elegans showed these to be recruited independently of each other and function redundantly in assembly of transition zone Y-links (Fig. 1 A; Williams et al., 2011). Proteomic analysis in vertebrates identified a potential third module including the key ciliopathy protein CEP290 (Fig. 1 B; Sang et al., 2011). Reciprocal BLAST searches identified a C. elegans homologue encoded by the predicted gene Y47G6A.17, which we named CCEP-290 for C. elegans CEP290 (Fig. S1, A and B). Phylegetic analysis found CEP290 to be conserved across all major eukaryotic phyla, supporting a central role for this protein at the transition zone (Fig. S1 C; Hodges et al., 2010).

Generation of a single-copy GFP transgene under control of endogenous regulatory sequences revealed expression exclusively in ciliated neurons in a manner dependent on the transcription factor DAF-19 (Fig. S1 D), which serves as a master regulator of ciliogenesis in C. elegans (Swoboda et al., 2000). Consistent with localization to transition zones, CCEP-290 was found distal to the basal body marker HYLS-1 (Dammermann et al., 2009) but proximal to the axoneme positive for poly-glutamyalted tubulin (Fig. 1 C). Comparisons by high-resolution two-color fluorescence imaging (Wan et al., 2009) using HYLS-1 as a point of reference showed this localization to be distinct from that of the MKS and NPHP module components MKS-2 and NPHP-4. Whereas MKS-2 and NPHP-4 localized to the transition zone periphery, consistent with localization to Y-links or the ciliary membrane, CCEP-290 occupied a more central position (Fig. S2, A and B). CCEP-290 recruitment was unaffected in mutants of MKS-5, MKS-2, and NPHP-4, the most upstream components of the MKS and NPHP assembly pathways (Fig. 1 D). CCEP-290 therefore independently targets to the transition zone.

Since C. elegans neurons are refractory to RNAi, we obtained an in-frame deletion mutant, ccep-290(tm4927), hereafter referred to as ccep-290, affecting the N terminus of the protein (Fig. S1 E). We also generated a complete gene deletion, ccep-290Δ, using CRISPR/Cas9-mediated genome editing (Fig. S1 F). Both mutants, as well as two additional C-terminal truncation mutants, were found to be viable and fertile, with no apparent morphological defects (Fig. S1 G and not depicted). Examining other transition zone proteins in ccep-290Δ mutants, we observed no change in localization of NPHP-4. In contrast, MKS-2 at transition zones appeared much reduced (Fig. 1 E), which is consistent with recent findings in Drosophila melanogaster (Basiri et al., 2014). Localization of downstream components of the MKS pathway was also perturbed (Fig. S2 C). Closer inspection revealed that MKS-2 still targeted to cilia (70.1 ± 16.0% of controls, t test, P < 0.0001). However, most of the protein (61.8 ± 16.4%) was now found dispersed along the axoneme, with MKS-2 particles moving in a manner characteristic of diffusion (Fig. 1 F and Video 1). Instead of targeting MKS-2 as expected for an upstream component in the MKS assembly pathway, CCEP-290 therefore tethers MKS-2 to the transition zone.

Examination of the C. elegans transition zone by serial-section electron microscopy reveals two prominent ultrastructural features: Y-links connecting axonemal microtubules to the ciliary membrane and the central cylinder to which axonemal outer doublet and inner singlet microtubules are attached (Fig. 1 G; Perkins et al., 1986). Tomographic reconstructions show both structures to be contiguous along the longitudinal axis of the transition zone, forming elongated sheets (Video 2, see also Fig. 4 A). Previous work in C. elegans had implicated MKS and NPHP proteins in assembly of Y-links. Loss of these connectors results in detachment and dilation of the ciliary membrane, while leaving the central cylinder and associated axonemal microtubules intact (Williams et al., 2011). The same role had been ascribed to CEP290 in Chlamydomonas reinhardtii, although Y-links were still observed, albeit rarely, in cep290 mutants (Craigie et al., 2010). In contrast, loss of CCEP-290 in C. elegans ccep-290Δ mutants resulted in fragmentation of the central cylinder, disrupting the radially symmetric array of axonemal microtubules (Fig. 1 G). Unlike in MKS/NPHP mutants, there was no detachment of the ciliary membrane and Y-links could still be found in some sections (arrowheads). ccep-290Δ;npbh-4 mutants displayed an intermediate phenotype, with 5/11 transition zones appearing fragmented and the remainder fully disorganized. Finally, disruption of all three modules in ccep-290; mksr-2; npbh-4 triple mutants resulted in a complete loss of transition zone structures, with axonemal microtubule doublets dissociated from each other and the ciliary membrane (Fig. 1 G). These results are consistent with our high-resolution localization data and highlight the distinct roles of CCEP-290 and MKS/NPHP proteins at the transition zone, with CCEP-290 serving as a core component of the central cylinder, which appears to act as an inner scaffold for transition zone assembly, while MKS/NPHP proteins function in assembly of peripheral Y-links.
Figure 2. Ciliogenesis is largely normal in transition zone mutants. (A) Phasmid cilia in wild-type and transition zone mutants visualized by the IFT marker CHE-11:GFP (arrowheads). (B and C) Quantitation of phasmid cilia number per animal (B, n > 50 animals) and length (C, n > 25 cilia) based on CHE-11:GFP. Error bars indicate the 95% confidence intervals. Asterisks indicate statistically significant difference to wild-type (t test, P < 0.01). (D) Stills and
We next examined the consequences of transition zone perturbations on cilia assembly. A commonly used method to assess cilia integrity in *C. elegans* is the dye-fill assay, which monitors uptake of the lipophilic dye Dil (Inglis et al., 2007). This dye is taken up by a subset of ciliated neurons (12 amphid neurons in the head, 4 phasmid neurons in the tail) through their exposed ciliated endings and accumulates in their cell bodies. Defective dye-filling (Dyf) is often indicative of compromised cilia structure. As previously reported for other transition zone mutants (Williams et al., 2008, 2011), *cccep-290* mutants individually did not display significant dye-fill defects. However, double mutants with either *nphp-1* or *nphp-4* were strongly Dyf. In contrast, no genetic interactions were detected with *mksr-2*. Dye-filling was restored by expression of the appropriate GFP transgene, confirming specificity of the mutant phenotypes and functionality of our GFP reporters (Fig. S1 H; Fig. S2, D and E; and not depicted). These results would appear to support a role for the transition zone in cilia assembly. However, direct visualization of phasmid cilia using the IFT marker CHE-11:GFP failed to reveal any major defects. Most strikingly, ~83% of cilia were found to remain in *cccep-290;mksr-2;*, *nphp-4* triple mutants in which all three transition zone modules are inhibited (Fig. 2, A and B). Cilia lengths (Fig. 2 C) and IFT rates (Fig. 2 D, E) were also largely normal. Ciliary ultrastructure was also unaffected with the exception of occasional displaced doublet microtubules, likely reflecting disorganization within the transition zone (Fig. 2 F). We conclude that loss of transition zone structures has only mild effects on axoneme assembly and organization in *C. elegans*.

While cilia assembly was largely unaffected in transition zone mutants, neuronal morphology was strongly perturbed. This was most clearly seen in phasmids, where dendrites collapsed almost entirely, with cilia found immediately adjacent to the cell body (Fig. 2 A). This phenomenon was previously reported in certain combinations of transition zone mutants (Williams et al., 2008, 2011). Loss of contact with the external environment rather than ciliogenesis defects could explain the observed lack of dye-filling, and dendrite lengths do correlate with dye-fill phenotypes (compare Figs. 3 C and S2 E). Complete dendrite collapse was not observed in amphids (Fig. S3, C and D). However, cilia frequently failed to extend into the channel formed by socket and sheath glia (Fig. S3, A and B).

Dendrite extension in amphids has been shown to occur by retrograde extension, whereby the cell body migrates backward while the dendritic tip remains anchored in place (Heiman and Shaham, 2009; Fig. 3, A and B; and Video 3). Failure of dendrite extension in transition zone mutants could reflect a failure of cell migration or dendrite anchorage. The stochastic nature of the phenotype and bilaterally symmetric organization of phasmids allowed us to distinguish between these possibilities. In cases where one set of phasmids displayed collapsed dendrites, their cell bodies were positioned opposite their counterparts with normally extended dendrites (Fig. 3 D), indicating that cell migration occurred normally while dendrite attachment was defective. Interestingly, dendrite lengths are bimodally distributed, either normal or fully collapsed (Fig. 3 C). Thus, attachment only occurs at the tip of the dendrite. If that attachment is lost, the dendrite collapses entirely.

Cilia could contribute to neuronal morphology in a variety of ways, including by signaling. However, dendrite extension occurs before cilia are fully formed (Sulston et al., 1983; Fujiwara et al., 1999) and was normal in IFT mutants in which axoneme assembly was inhibited (Snow et al., 2004; Fig. S3 H). Because signaling requires an intact cilium, this would support a direct anchoring role rather than an indirect signaling one, mediated by some structure at the ciliary base. A unique feature of *C. elegans* is that centrioles degenerate early during ciliogenesis (Dammernmann et al., 2009) and do not persist at the base of mature cilia (Perkins et al., 1986). While transition zone proteins are present at the time of dendrite elongation (Fig. 3 E), centriole degeneration is well underway at this time (Fig. S3 I), making it unlikely that centrioles or centrosomes contribute to dendrite anchorage. Finally, loss of the basal body protein HYLS-1 only mildly perturbed dendrite extension in a manner synergistic with loss of NPHP-4 (Fig. S3 H), likely reflecting transition zone disorganization in *hyls-1* mutants. Our results are therefore most consistent with a direct anchoring function mediated by the transition zone.

Dendrite extension in PQR neurons has been reported to occur in the canonical, anterograde manner by growth cone crawling (Kirszenblat et al., 2011). It is therefore not surprising that dendrites in PQR neurons are unaffected (Fig. S3, E and F). In contrast, both amphid and phasmid dendrites form by retrograde extension. Why the more severe phenotype in phasmids? We speculate that this is due to cooperativity in attachment between neurons sharing the same channel. Indeed, dendrites of the two neurons sharing each phasmid channel always succeeded or failed to extend together, whereas there was no correlation between dendrite extension on the left and right sides of the animal (65% displayed collapsed dendrites on the left; 63% on the right; 41% both sides, the expected probability for two independent events; n = 46 animals). With 13 transition zones per amphid compared with two per phasmid, cooperative attachment could make amphid dendrites more resistant to collapse. Differences in cell morphology may also contribute. It is notable in this context that failure of dendrite extension in phasmids, unlike in amphids (Heiman and Shaham, 2009), is not accompanied by failure of sheath glia extension (Fig. S3 G).

For a structure at the dendritic tip to provide anchorage during retrograde extension, it must interact with neighboring cells or other immobile structures. For amphids, that structure is the extracellular matrix, composed in part of the tectorin-related proteins DEX-1 and DYF-7. Loss of either protein results in loss of anchorage and dendrite collapse (Heiman and Shaham, 2009). This was also true for the strong loss-of-function mutant *dyf-7(m537)* in phasmids (Fig. 3 F), indicating that dendrites are secured to the extracellular material as in amphids. This adhesive matrix remains in place in transition zone mutants (Fig. 3 G).
Figure 3. **A role for the transition zone in dendrite attachment.** (A) Schematic of dendrite formation by retrograde extension in *C. elegans* amphids. (B) Stills from time-lapse movie of embryo expressing myristoylated GFP in amphid neurons undergoing retrograde migration. Overlay of GFP signal is shown on transmitted light images of the embryo. Insets show GFP only. Note that the position of dendritic tips (arrowhead) appears fixed as cell bodies...
If transition zones and tectorins function together in cell adhesion, we should be able to observe genetic interactions between mutants in both structures. Indeed, hypomorphic mutants of *dex-1* and *dyf-7* were found to synergize with mutations in *nphp-4*, which alone do not result in dendrite phenotypes (Fig. 3, C and F). Thus, transition zones mediate dendrite attachment via interactions with the extracellular matrix.

In summary, our work has helped clarify the molecular architecture and function of the ciliary transition zone (see also Fig. 4, A and B). By combining perturbation of CCEP-290, which affects the central cylinder, with previously described perturbations that affect the Y-links (Williams et al., 2011), we were able to completely disrupt transition zone structure. Our finding, that axoneme assembly remains relatively normal when the transition zone is lost, argues that there is no fundamental requirement for the transition zone in axoneme assembly. The picture has been less clear in vertebrates, where loss of transition zone components has been reported to result in ciliary defects in some tissues but not others (Garcia-Gonzalo et al., 2011; Sang et al., 2011; Chih et al., 2012). One possible explanation for this context dependence is that it relates to transition zone function in ciliary gating (Craigie et al., 2010). Perturbed entry of ciliary components could affect cilia assembly in some cases but not others. Alternatively, loss or impairment of transition zone structures could affect cilia stability. As previously shown for basal bodies in *Tetrahymena thermophila* (Bayless et al., 2012), cilia motility or fluid flow over nonmotile cilia results in mechanical stresses that could damage cilia unless stabilized by structures at their base. In less challenging environments, such as *C. elegans* cilia protected inside the cuticle of the worm, impaired stability might have less deleterious consequences.

A second major conclusion of our work is that the transition zone is required for adhesion to the extracellular matrix to allow dendrite extension. While cilia have not previously been implicated in cell adhesion, signs of crosstalk have been observed for many years. Thus, NPHP proteins localize to cell–cell and cell–matrix junctions (Benzing et al., 2001) and have been found to be required for epithelial morphogenesis (Delous et al., 2009). Conversely, junctional proteins have been found at cilia (Fan et al., 2004; Sfakianos et al., 2007). Cilia have been linked to many developmental processes in vertebrates and we speculate that cilia-mediated adhesion contributes to some of these processes. For example, during development of the cerebral cortex, delamination of neural precursors from the apical adherens belt is an important determinant of cell fate. These precursors are ciliated, and it has been shown that the position of the primary cilium determines whether a daughter cell delaminates after mitosis (Wilsch-Bräuninger et al., 2012; Paridaen et al., 2013; Fig. 4 C). Importantly, cilia repositioning precedes any change in polarity and is therefore the driving force behind loss of attachment. While it is possible that cilia-dependent signaling underlies this phenomenon, we suggest an alternative explanation: a direct, mechanical role for the cilium in cell adhesion. As in *C. elegans*, analysis of IFT mutants (impairing cilia-dependent signaling but not transition zone function) could distinguish between these possibilities.

**Materials and methods**

**Sequence analysis**

CCEP290 orthologues were identified by reciprocal BLAST analysis using human CCEP290 as the starting point. Where direct comparisons failed to identify a clear homologue, indirect searches were performed using less divergent related species as intermediates. Orthology relationships were confirmed using an algorithm optimized for coiled-coil proteins (Kuhn et al., 2014). Conserved motifs within CCEP290 were identified by Meme (http://meme-suite.org/) and sequences were aligned using Mafft within Jalview (http://www.jalview.org). Maximum likelihood phylogenetic trees were constructed using IQ-TREE under the LG+G model (Le and Gascuel, 2008). Clade supports were assessed as the ultrafast bootstrap (UFBoot) analysis (Minh et al., 2013).

**C. elegans strains and culture conditions**

Strains used are listed in Table S1. *ccep-290* mutants were obtained from the *C. elegans* Knockout Consortium/NBRP-Japan (m4927) and the Million Mutation Project (gk415029 and gk107339). The *ccep-290Δ* mutant was generated by Cas9-induced homologous recombination (Dickinson et al., 2013), replacing the entire *ccep-290* coding region and 5′ and 3′ UTR with the selectable marker unc-119. Detailed information for other mutants is available from Wormbase (http://www.wormbase.org). Mutants were outcrossed 6× to N2 wild-type before phenotypic analysis and introduction of fluorescent markers. Transgenic strains expressing fluorescently tagged CCEP-290, HYLS-1, MKSR-2, and NPHP-4 were generated by cloning the corresponding genomic loci, including 5′ and 3′ regulatory sequences and N-terminal LAP-GFP/mCherry into the appropriate targeting vector and Mos1-mediated transposition (Frøkjaer-Jensen et al., 2008). Strains expressing GFP:CCEP-290, DYF-7:GFP, MKS-2:GFP, MKS-6:GFP, MKSR-1:GFP, and Pnas-31:mCherry/OSM-6:GFP from extrachromosomal arrays were generated by coinjection with the marker plasmid pRF4. All strains were maintained at 23°C.

**Immunofluorescence and fixed imaging**

Immunofluorescence experiments were performed as described previously (Oegema et al., 2001) using Cy3/Cy5-labeled affinity-puriﬁed rabbit polyclonal antibodies against HYLS-1 (Dammermann et al., 2009) and SAS-4 (Dammermann et al., 2004), as well as unlabeled mouse monoclonal antibodies against polyglutamylated tubulin (GT335; provided by C. Janke, Institut Curie, Paris, France). Residual (asterisk) move. See also Video 3. (C) Quantitation of phasmdid dendrite lengths in wild-type and transition zone mutants at the L4 stage, using CHE-11:GFP. Each dot on the scatter plot represents a single dendrite, with bars indicating average and 95% conﬁdence intervals. n > 20 dendrites per condition. Asterisks indicate statistically signiﬁcant differences to wild-type (*t* test; ***, *P < 0.001*). (D) Tail of a *ccep-290; nphp-1* mutant animal expressing CHE-11:GFP. While dendrite extension has failed in both phasmids on the left hand side, right-hand-side phasmids have formed normally. Note that cell bodies are positioned similarly on both sides (broken line). (E) Immunofluorescence micrographs of amphids and phasmds of late embryos expressing Myc-stained GFP and LAP-CCEP-290 stained for CCEP-290 with antibody to S-tag. CCEP-290 is located at the tip of the elongating dendrite (arrowheads). (F) Phasmdid dendrite lengths in combinations of *dex-1*/*dyf-7* and *nphp-4* mutants, quantitated as described in comparison. ns42 and ns117 are hypomorphic alleles of *dex-1* and *dyf-7*, *dyf-7(m537)* closer to a loss of function (Heiman and Shaham, 2009). Asterisks indicate statistically signiﬁcant differences to *dex-1*/*dyf-7* single mutants (*t* test; ***, *P < 0.001*; **, *P < 0.01*). (G) Still images of wild-type and *ccep-290; nphp-4* mutant embryos expressing DVF-7:GFP. Overlay of GFP signal on transmitted light images of the embryo. Note the enrichment of DVF-7:GFP at the tips of elongating dendrites before phenotypic analysis and introduction of fluorescent markers.
fluorescence signal was used to detect GFP/mCherry-tagged protein in fixed embryos. Mouse monoclonal antibodies against S-peptide (MA1-981; Thermo Fisher Scientific) were used to detect LAP-tagged CCEP-290 in embryos also expressing other GFP fusions. In brief, worms were rehydrated by freeze crack, fixed in −20°C methanol for 20 min, rehydrated in PBS, blocked for 20 min in AbDil (PBS, 2% BSA, 0.1% Triton X-100), incubated with primary antibodies (1 µg/ml HYLS-1/SAS-4/MA1-981, 4 µg/ml GT335) in AbDil for 1.5 h, washed with PBST (PBS, 0.1% Triton X-100), incubated with secondary antibodies (15 µg/ml Cy2/Cy3/Cy5-labeled donkey anti-mouse antibody; Jackson ImmunoResearch Laboratories, Inc.) in AbDil for 1.5 h, washed with PBST, and incubated with 1 µg/ml Hoechst 33342 in PBS for 5 min before mounting in 0.5% p-phenylenediamine, 20 mM Tris, pH 8.8, and 90% glycerol. 3D wide-field datasets were acquired using a 100x 1.4 NA Super Plan-Apochromat lens on a microscope (DeltaVision; Applied Precision) equipped with a 7-Color SS1 module and a cooled charge-coupled device (CCD) camera (CoolSNAP-HQ2; Photometrics), computationally deconvolved using the enhanced ratio constrained iterative deconvolution algorithm, and maximum-intensity projection in SoftWorx (Applied Precision), before being imported into Photoshop (Adobe) for panel preparation. No nonlinear gamma corrections were performed during image processing.

**Live imaging**
Where possible, worms expressing fluorescent markers were examined without fixation, anesthetized with 10 mM tetramisole, and mounted on 2% agarose pads. Imaging was performed at room temperature on the set-up described above. For examination of neuronal morphology and localization interdependencies, 3D wide-field datasets were acquired, deconvolved, and processed as described above. Exposure conditions and scaling were kept constant when comparing signal across mutant backgrounds. At least 10 animals were examined per condition.

Figure 4. **Molecular architecture of the transition zone and function in cell adhesion.** (A) 3D reconstruction model of the transition zone in *C. elegans*, highlighting key features. See also Video 2. Based on protein localization and mutant phenotypes, CCEP-290 is an essential component of the central cylinder (this study), whereas MKS and NPHP module components function in assembly of Ylinks (Williams et al., 2011). Transition zone mutants display no apparent defects in axoneme assembly. However, ciliary gating and cell adhesion are compromised. (B) The dendritic of *C. elegans* amphid and phasmid neurons forms by retrograde extension, with the cell body moving backward while the dendritic tip remains in place. The transition zone mediates tip anchorage via interactions with the extracellular matrix. (C) Positioning of the primary cilium determines cell fate in vertebrate neuroepithelium. Cells with apically positioned cilia maintain their position at the apical adherens junction belt, whereas cells with basolateral cilial delaminate (Wilsch-Bräuninger et al., 2012; Paridaen et al., 2013). Differential signaling, but also mechanical anchorage by the cilium, could explain this result.
a CCD camera (Eagle 4k HS; FEI) and operated at 200 kV. Single axis tilt series were acquired using SerialEM (Mastronarde, 2005) by stepwise tilting the sample on average from −60° to +60° in 1° increments. Tomogram reconstruction with 10-nm gold beads as fiducials and model generation was performed using the IMOD software package (Kremer et al., 1996).

Online supplemental material

Fig. S1 details the identification and initial characterization of C. elegans CEP290, including sequence alignments, expression analysis, and phenotypic characterization of ccep-290 mutants. Fig. S2 presents a high-resolution analysis of transition zone protein localization and localization interdependencies, as well as an analysis of genetic interactions between transition zone mutants as revealed by the dye-fill assay. Fig. S3 presents a further characterization of dendrite extension, including phenotypes in other ciliated neurons, glial cell morphology, and contribution of other ciliary structures. Table S1 lists C. elegans strains used in this study. Table S2 provides accession numbers of CEP290 orthologues. Video 1 shows MKSR-2 particle movement along the ciliary axoneme in a ccep-290 mutant. Video 2 shows an electron tomogram and 3D reconstruction model of the C. elegans transition zone. Video 3 shows dendrite extension by retrograde migration in the C. elegans embryo. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201501013/DC1.

Acknowledgements

We thank Karen Oegema for comments on the manuscript, the Vienna Biocenter C. elegans community, and members of the Dammermann laboratory for discussions; the Caenorhabditis Genetics Center, Max Heiman (Boston Children’s Hospital, Boston, MA), Carsten Janke, Michel Leroux (Simon Fraser University, Burnaby BC, Canada), Shohei Mitani (Tokyo Women’s Medical University, Tokyo, Japan), and Bradley Yoder (The University of Alabama at Birmingham, Birmingham, AL) for strains and reagents, Bui Quang Minh for help with bioinformatics, and the Campus Science Support Facilities GmbH (CSF) EM facility for technical assistance.

This work was supported by startup funding from Max F. Perutz Laboratories and grants Y597820 and P24296820 from the Austrian Science Fund (FWF) to A. Dammermann, as well as a predoctoral fellowship from the Boehringer Ingelheim Fonds to C. Schouteden.

The authors declare no competing financial interests.

Submitted: 5 January 2015
Accepted: 1 June 2015

References

Basir, M.L., A. Ha, A. Chadha, N.M. Clark, A. Polyanyovsky, B. Cook, and T. Avidor-Reiss. 2014. A migrating cilia gate compartmentalizes the site of axoneme assembly in Drosophila spermatids. Curr. Biol. 24:2622–2631. http://dx.doi.org/10.1016/j.cub.2014.09.047

Bayless, B.A., T.H. Giddings Jr., M. Winey, and C.G. Pearson. 2012. Bld10/Chlamydosomus kinesin-II-dependent intraflagellar transport (IFT). IFT particles contain proteins required for ciliary assembly in Caenorhabditis elegans sensory neurons. J. Cell Biol. 141:993–1008. http://dx.doi.org/10.1083/jcb.141.4.993

Craig, B., C.C. Tsao, D.R. Diener, Y. Hou, F.K. Lechtreck, J.L. Rosenbaum, and G.B. Wittman. 2010. CEP290 tethers flagellar transition zone microtubules to the membrane and regulates flagellar protein content. J. Cell Biol. 190:927–940. http://dx.doi.org/10.1083/jcb.201006105

Dammermann, A., T. Müller-Reichert, L. Pelletier, B. Habermann, A. Desai, and K. Oegema. 2004. Centriole assembly requires both centriolar and pericentriolar materials. Dev. Cell. 7:815–829. http://dx.doi.org/10.1016/j.devcel.2004.10.015

Dammermann, A., H. Pemble, B.J. Mitchell, I. McLeod, J.R. Yates III, C. Kintner, A.B. Desai, and K. Oegema. 2009. The hydroxyethyl sulfonamide protein HYLS-1 links core centriole structure to cilia formation. Genes Dev. 23:2046–2059. http://dx.doi.org/10.1101/gad.1810409

Delous, M., N.E. Hellman, H.M. Gaude, F. Silbernagl, A. Le Bivic, R. Salomon, C. Antignac, and S. Saumier. 2009. Nephrocystin-1 and nephrocystin-4 are required for epithelial morphogenesis and associate with PALS1/PATJ and Par6. Hum. Mol. Genet. 18:4711–4723. http://dx.doi.org/10.1093/hmg/ddp434

Dickinson, D.J., J.D. Ward, D.J. Reiner, and B. Goldstein. 2013. Engineering the Caenorhabditis elegans genome using Cas9-triggered homologous recombination. Nat. Methods. 10:1026–1034. http://dx.doi.org/10.1038/nmeth.2641

Fan, S., T.W. Hurd, C.J. Liu, S.W. Straight, T. Weimbs, E.A. Hurd, S.E. Domino, and B. Margolis. 2004. Polarity proteins control ciliogenesis via kinesin motor interactions. Curr. Biol. 14:1451–1461. http://dx.doi.org/10.1016/j.cub.2004.08.025

Fliiegauf, M., T. Benzing, and H. Omran. 2007. When cilia go bad: cilia defects and ciliopathies. Nat. Rev. Mol. Cell Biol. 8:880–893. http://dx.doi.org/10.1038/nrm2278

Frukjaer-Jensen, C., M.W. Davis, C.E. Hopkins, B.J. Newman, J.M. Thummel, S.P. Olesen, M. Grunnet, and E.M. Jorgensen. 2008. Single-copy insertion of transgenes in Caenorhabditis elegans. Nat. Genet. 40:1375–1383. http://dx.doi.org/10.1038/ng.248

Fujiwara, M., T. Ishihara, and I. Katsura. 1999. A novel WD40 protein, CHE-2, acts cell-autonomously in the formation of C. elegans sensory cilia. Development. 126:4839–4848.

Garcia-Gonzalo, F.R., K.C. Corbit, M.S. Sirerol-Piquer, G. Ramaswami, E.A. Otto, T.R. Noriega, A.D. Seel, J.F. Robinson, Č.L. Bennett, D.J. Josifova, et al. 2011. A transition zone complex regulates mammalian ciliogenesis and ciliary membrane composition. Nat. Genet. 43:776–784. http://dx.doi.org/10.1038/ng.891

Heiman, M.G., and S. Shaham. 2009. DEX-1 and DYE-7 establish sensory dendrite length by anchoring dendritic tips during cell migration. Cell. 137:344–355. http://dx.doi.org/10.1016/j.cell.2009.01.057

Hodges, M.E., N. Scheumann, B. Wickstead, J.A. Langdale, and K. Gull. 2010. Reconstructing the evolutionary history of the centriole from protein components. J. Cell Sci. 123:1407–1413. http://dx.doi.org/10.1242/jcs.068873

Huang, L., K. Szymanska, V.L. Jensen, A.R. Janecke, A.M. Innes, E.E. Davis, P. Frosk, C. Li, J.R. Willer, B.N. Chodirker, et al. 2011. TMEM237 is mutated in individuals with a Joubert syndrome related disorder and expands the role of the TMEM family at the ciliary transition zone. Am. J. Hum. Genet. 89:793–730. http://dx.doi.org/10.1016/j.ajhg.2011.11.005

Inglis, P.N., G. Ou, M.R. Leroux, and J.M. Scholey. 2007. The sensory cilia of Caenorhabditis elegans (March 8, 2007). In WormBook. The C. elegans Research Community, Editors. http://dx.doi.org/10.1895/ wormbook.1.126.2

Kirschstal, L., D. Pattabiraman, and M.A. Hilliard. 2011. LIN-44/Wnt directs dendrite outgrowth through LIN-17/Prized in C. elegans neurons. PLoS Biol. 9:e1001157. http://dx.doi.org/10.1371/journal.pbio.1001157

Kremer, J.R., D.N. Mastronarde, and J.R. McIntosh. 1996. Computer visualization of three-dimensional image data using IMOD. J. Struct. Biol. 116:71–76. http://dx.doi.org/10.1006/jsbi.1996.0013

Kuhn, M., A.A. Hyman, and A. Beyer. 2014. Coiled-coil proteins facilitated the functional expansion of the centrosome. PLoS Comput. Biol. 10:e1003657. http://dx.doi.org/10.1371/journal.pcbi.1003657

Le, S.Q., and O. Gascuel. 2008. An improved general amino acid replacement matrix. Mol. Biol. Evol. 25:1307–1320. http://dx.doi.org/10.1093/molbev/msn067

Mastronarde, D.N. 2005. Automated electron microscopy tomography using robust prediction of specimen movements. J. Struct. Biol. 152:36–51. http://dx.doi.org/10.1016/j.jsb.2005.07.007

Ciliary transition zone function in C. elegans • Schouteden et al. 43
Minh, B.Q., M.A. Nguyen, and A. von Haeseler. 2013. Ultrafast approximation for phylogenetic bootstrap. Mol. Biol. Evol. 30:1188–1195. http://dx.doi.org/10.1093/molbev/mst024

Oegema, K., A. Desai, S. Rybina, M. Kirkham, and A.A. Hyman. 2001. Functional analysis of kinetochore assembly in Caenorhabditis elegans. J. Cell Biol. 153:1209–1226. http://dx.doi.org/10.1083/jcb.153.6.1209

Paridaen, J.T., M. Wilsch-Bräuninger, and W.B. Huttner. 2013. Asymmetric inheritance of centrosome-associated primary cilium membrane directs ciliogenesis after cell division. Cell. 155:333–344. http://dx.doi.org/10.1016/j.cell.2013.08.060

Perkins, L.A., E.M. Hedgecock, J.N. Thomson, and J.G. Culotti. 1986. Mutant sensory cilia in the nematode Caenorhabditis elegans. Dev. Biol. 117:456–487. http://dx.doi.org/10.1016/0012-1606(86)90314-3

Reiter, J.F., O.E. Blacque, and M.R. Leroux. 2012. The base of the cilium: roles for transition fibres and the transition zone in ciliary formation, maintenance and compartmentalization. EMBO Rep. 13:608–618. http://dx.doi.org/10.1038/embor.2012.73

Rosenbaum, J.L., and G.B. Witman. 2002. Intraflagellar transport. Nat. Rev. Mol. Cell Biol. 3:813–825. http://dx.doi.org/10.1038/nrm952

Sang, I., J.J. Miller, K.C. Corbit, R.H. Giles, M.J. Brauer, E.A. Otto, L.M. Baye, X. Wen, S.J. Scales, M. Kwong, et al. 2011. Mapping the NPHP-JBTS-MKS protein network reveals ciliopathy disease genes and pathways. Cell. 145:513–528.

Schmidt, K.N., S. Kuhns, A. Neuner, B. Hub, H. Zentgraf, and G. Pereira. 2012. Cep164 mediates vesicular docking to the mother centriole during early steps of ciliogenesis. J. Cell Biol. 199:1083–1101. http://dx.doi.org/10.1083/jcb.201202126

Sfakianos, J., A. Togawa, S. Maday, M. Hull, M. Pypaert, L. Cantley, D. Toomre, and I. Mellman. 2007. Par3 functions in the biogenesis of the primary cilium in polarized epithelial cells. J. Cell Biol. 179:1133–1140. http://dx.doi.org/10.1083/jcb.200709111

Snow, J.J., G. Ou, A.L. Gunnarson, M.R. Walker, H.M. Zhou, I. Brust-Mascher, and J.M. Scholey. 2004. Two anterograde intraflagellar transport motors cooperate to build sensory cilia on C. elegans neurons. Nat. Cell Biol. 6:1109–1113. http://dx.doi.org/10.1038/ncb1186

Sorokin, S. 1962. Centrioles and the formation of rudimentary cilia by fibroblasts and smooth muscle cells. J. Cell Biol. 15:364–377. http://dx.doi.org/10.1083/jcb.15.2.364

Sulston, J.E., E. Schierenberg, J.G. White, and J.N. Thomson. 1983. The embryonic cell lineage of the nematode Caenorhabditis elegans. Dev. Biol. 100:64–119. http://dx.doi.org/10.1016/0012-1606(83)90201-4

Swoboda, P., H.T. Adler, and J.H. Thomas. 2000. The RFX-type transcription factor DAF-19 regulates sensory neuron cilium formation in C. elegans. Mol. Cell. 5:411–421. http://dx.doi.org/10.1016/S1097-2765(00)80436-0

Wan, X., R.P. O’Quinn, H.L. Pierce, A.P. Joglekar, W.E. Gall, J.G. DeLuca, C.W. Carroll, S.T. Liu, T.J. Yen, B.F. McEwen, et al. 2009. Protein architecture of the human kinetochore microtubule attachment site. Cell. 137:672–684. http://dx.doi.org/10.1016/j.cell.2009.03.035

Williams, C.L., M.E. Winkelbauer, J.C. Schafer, E.J. Michaud, and B.K. Yoder. 2008. Functional redundancy of the B9 proteins and nephrocystins in Caenorhabditis elegans ciliogenesis. Mol. Biol. Cell. 19:2154–2168. http://dx.doi.org/10.1091/mbc.E07-10-1070

Williams, C.L., C. Li, K. Kida, P.N. Inglis, S. Mohan, L. Semenec, N.J. Bialas, R.M. Stupay, N. Chen, O.E. Blacque, et al. 2011. MKS and NPHP modules cooperate to establish basal body/transition zone membrane associations and ciliary gate function during ciliogenesis. J. Cell Biol. 192:1023–1041. http://dx.doi.org/10.1083/jcb.201012116

Wilsch-Bräuninger, M., J. Peters, J.T. Paridaen, and W.B. Huttner. 2012. Basolateral rather than apical primary cilium on neuroepithelial cells committed to delamination. Development. 139:95–105. http://dx.doi.org/10.1242/dev.062949