The small GTPases ARL-13 and ARL-3 coordinate intraflagellar transport and ciliogenesis

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Intraflagellar transport (IFT) machinery mediates the bidirectional movement of cargos that are required for the assembly and maintenance of cilia. However, little is known about how IFT is regulated in vivo. In this study, we show that the small guanosine triphosphatase (GTPase) adenosine diphosphate ribosylation factor–like protein 13 (ARL-13) encoded by the Caenorhabditis elegans homologue of the human Joubert syndrome causal gene ARL13B, localizes exclusively to the doublet segment of the cilium. arl-13 mutants have shortened cilia with various ultrastructural deformities and a disrupted association between IFT subcomplexes A and B. Intriguingly, depletion of ARL-3, another ciliary small GTPase, partially suppresses ciliogenesis defects in arl-13 mutants by indirectly restoring binding between IFT subcomplexes A and B. Rescue of arl-13 mutants by ARL-3 depletion is mediated by an HDAC6 deacetylase-dependent pathway. Thus, we propose that two conserved small GTPases, ARL-13 and ARL-3, coordinate to regulate IFT and that perturbing this balance results in cilia deformation.

Introduction

Cilia, which act as motile or sensory devices on the surfaces of most eukaryotic cells, play essential roles in development and postnatal tissue pattern formation. Mounting evidence links defects in cilia biogenesis and/or function to a wide spectrum of human genetic disorders that have been collectively termed ciliopathies (Badano et al., 2006). Joubert syndrome (JS) is an autosomal recessive disorder characterized by abnormalities in the central nervous system and by various commonly shared ciliopathy manifestations such as cystic kidney, blindness, and polydactyly. The functions of all identified JS loci (NPHP1, AHI1, CEP290, RPGRIP1L, TMEM67/MKS3, Arl13B, and CC2D2A), as well as most other ciliopathy candidate genes, have been suggested to be cilia and/or basal body related. Nonetheless, the precise roles of these genes remain enigmatic (Pazour and Rosenbaum, 2002; for reviews see Eley et al., 2005; Fliegauf et al., 2007; Parisi, 2009).

Intraflagellar transport (IFT) builds and maintains all cilia and flagella (Rosenbaum and Witman, 2002). In Caenorhabditis elegans, the head amphid and tail phasmid sensory organs terminate with channel cilia, which are composed of middle segments containing nine peripheral doublet microtubules and distal segments containing singlet microtubules (Perkins et al., 1986). In C. elegans, IFT is regulated by two anterograde motor machineries, the canonical heterotrimeric kinesin-II and homodimeric OSM-3/KIF17, and one retrograde dynein motor (Scholey, 2008). The two kinesin motors cooperate to build the middle doublets, whereas OSM-3 acts alone to extend distal singlets (Ou et al., 2005). Kinesin-II associates with IFT subcomplex A (IFT-A), OSM-3 associates with IFT subcomplex B (IFT-B), and the Bardet–Biedl syndrome (BBS) proteins stabilize the association between IFT-A and IFT-B. In the absence of BBS-7 and BBS-8, IFT-A and IFT-B separate as a result of mechanical competition between the two kinesin motors (Ou et al., 2005). Remarkably, both kinesin-II/KIF3 and OSM-3/KIF17 homologues appear to contribute to IFT and ciliogenesis in vertebrate systems (Marszalek et al., 2000; Lin et al., 2003; Jenkins et al., 2006; Insinna et al., 2008, 2009).

ADP ribosylation factor (Arf)–like proteins (Arls) are enzymatically active small GTPases in the Ras superfamily. The family of mammalian Arf/Arl GTPases includes 29 members, and the roles of most Arf/Arls are poorly understood (for original data in the JCB Data Viewer http://jcb-dataviewer.rupress.org/jcb/browse/2862
reviews see D’Souza-Schorey and Chavrier, 2006; Gillingham and Munro, 2007). In comparative genomic searches, ARL3 and Arl6 were identified as genes that are expressed exclusively in ciliated organisms (Avidor-Reiss et al., 2004; Li et al., 2004). Later, human Arl6 was identified as a locus causative for BBS syndrome, and its C. elegans homologue was found to enter the cilium and undergo IFT (Fan et al., 2004). Meanwhile, ARL-3 was found in the Chlamydomonas reinhardtii ciliary proteome (Pazour et al., 2005) and implicated in Leishmania donovani flagellum biogenesis (Cuvillier et al., 2000). The Arn3<sup>−/−</sup> mouse dies after birth and exhibits typical ciliopathy manifestations such as cystic kidney and photoreceptor degeneration (Schrick et al., 2006). In addition, mutations in another small GTPase, ARL13b, leads to a ciliogenesis defect in which the C12–15 palmitoylation motif has been deleted, localized to both cilia and the cytosol (Fig. S1 D). Quantitation of fluorescence intensity revealed that only ~30% of the ARL-13<sup>C12-15Del</sup> protein localized to cilia compared with ~95% of wild-type ARL-13. This observation suggests that ARL-13 is a ciliary membrane-binding small GTPase.

C-terminal PRD restricts ARL-13 to the doublet segment of the cilium

In contrast to most small GTPases, which are ~20 kD and composed of only a GTPase domain, ARL13B has an atypical long C terminus that contains a coiled-coil helix and a PRD (Fig. 1 A). This unique structural pattern is conserved in all ARL13B protein family members from the green algae C. reinhardtii to mammals. In zebrafish, ciliary targeting of ARL13B requires both the N and C termini (Duldulao et al., 2009). To characterize the putative cilia targeting signal sequence of C. elegans ARL-13, we generated transgenic animals that express different GFP-tagged ARL-13 truncations. Interestingly, both the ARL-13 NT truncation (which retains the GTPase domain plus the coiled-coiled helix) and the ARL-13 PRD truncation (which possesses only the PRD domain) were targeted to cilia (Fig. 1 C), indicating redundant ciliary targeting signals in the N and C termini of the worm ARL-13 protein. Nonetheless, ARL-13 PRD::GFP showed restricted middle segment localization, whereas ARL-13 NT::GFP was present in both middle and distal segments. This result suggests that the PRD domain is critical for restriction of ARL-13 to the doublet segment. Although full-length ARL-13 protein was completely absent from the cell body, we observed significant GFP signals in the cytoplasmic region for ARL-13 NT::GFP and ARL-13 PRD::GFP (Fig. 1 C), which suggests that both the N and C termini are required for efficient ciliary targeting of ARL-13.

ARL-13 is required for amphid and phasmid ciliogenesis

Two deletion alleles are available for the <i>arl-13</i> gene, <i>gk513</i> and <i>tm2322</i>. The <i>gk513</i> allele removes a 1.1-kb DNA fragment, including the promoter and first exon (Fig. 1 A). The <i>tm2322</i> allele, an in-frame deletion that removes exons 7–9, encodes a truncated protein with an intact GTPase domain (Fig. 1 A). Both <i>arl-13(gk513)</i> and <i>arl-13(tm2322)</i> animals are healthy and show no gross anatomical abnormalities. We used <i>gk513</i> as a reference allele. Because of the unique expression pattern of the ARL-13 protein (Fig. 1 B), we focused our experiments on cilia.
In *C. elegans*, cilia integrity can be assessed by a simple and efficient dye-filling assay. In living worms, amphid and phasmid neurons take up lipophilic fluorescent dye through sensory cilia that are exposed to the outside environment (Hedgecock et al., 1985). Worm mutants with abnormal cilia cannot take up dye and are termed dye-filling defective (Dyf; Perkins et al., 1986). Under the fluorescence dissecting microscope, *arl-13 (gk513)* amphids showed normal dye uptake, whereas ~40% of phasmids were Dyf (Fig. 2 A). Although no Dyf phenotype was observed in amphids, our transmission electron microscopy (TEM) experiments revealed that some *arl-13* amphid cilia were malformed (see Fig. 4). The Dyf phenotype of *arl-13* mutants could be fully rescued by introducing an ARL-13::GFP transgene (Fig. 2 A). Furthermore, overexpression of ARL-13...
partially rescue the arl-13 (gk513) Dyf phenotype (Fig. 2 A). Overexpression of ARL-13R83Q also caused Dyf in wild-type animals. Collectively, our data suggest a conserved role for the ARL-13 protein and its GTPase activity in ciliogenesis.

arl-13 mutants show various cilia deformations and compromised IFT

We used ciliary markers to examine cilia morphology in arl-13 mutants. In arl-13 mutants, the IFT-B component CHE-11::GFP and the kinesin-I motor KAP-1::GFP showed no accumulation along cilia, whereas IFT-A components (CHE-13::GFP and OSM-5::GFP), the IFT-B-associated motor OSM-3, and BBS proteins (BBS-7::GFP and BBS-8::GFP) aggregated in similar bulblike structures. GTP-tagged CHE-11, KAP-1, OSM-6, and OSM-3 were expressed in wild-type and arl-13 worms, and IFT transport event ratios within middle segments of phasmid cilia were quantified. Note that the curves of the IFT-A component CHE-11 and kinesin-I subunit KAP-1 significantly shifted to the left for arl-13 mutants compared with wild-type worms, suggesting that significant amounts of IFT-A and the associated kinesin-I move at a slower rate in the mutants. In contrast, a right shift of the curves for the IFT-B component OSM-6 and the OSM-3 motor was observed, indicating that IFT-B and the associated OSM-3 motor tend to move at a faster rate in ARL-13–deficient cilia. In all figures, solid arrows indicate the TZs, dashed arrows point to the approximate middle to distal segment transition points, and arrowheads mark the distal ends. Statistical analysis was performed using a two-tailed paired Student’s t-test. *, P < 0.01. Error bars indicate SEM. Bars, 5 μm.
markers, we introduced each gfp transgene from wild-type strains into arl-13 mutants by genetic crossing. Expression of OSM-6::GFP, an IFT-B marker, revealed that ~40% phasmid cilia are truncated, and the cilia often show abnormal bulblike structures in the middle segments (Fig. 2 C). This ratio is comparable with the Dyf ratio of arl-13(gk513) phasmid cilia.

Further analyses revealed that IFT-B proteins (OSM-5, OSM-6, and CHE-13), the IFT-B–associated OSM-3 motor, and IFT regulators (BBS-7 and BBS-8) all accumulated in bulblike structures in middle segments, whereas the IFT-A protein CHE-11 and its associated motor KAP-1 did not exhibit bulblike accumulation along the cilium (Fig. 2 D). All observed bulblike structures were located proximal to the middle to distal transition point (Fig. 2 D, dashed arrows), which might indicate a defective transition of IFT-B subcomplexes (including associated BBS proteins and the OSM-3 motor) from middle segments to distal segments in arl-13 mutants. In transition zones (TZs; Fig. 2 D, solid arrows), all IFT markers showed similar enrichment in both arl-13 and wild-type worms.

Next, we examined IFT motility in arl-13 mutants. In wild-type animals, kinesin-II and OSM-3 cooperate in moving the same IFT particle along the middle segment at 0.7 µm/s. Then, the OSM-3 kinesin alone moves the IFT particle along the distal segment at the faster rate of 1.3 µm/s. Cytoplasmic dynein mediates retrograde movement of the IFT particle along the whole axoneme at a rate of 1.1 µm/s (Signor et al., 1999; Snow et al., 2004; Ou et al., 2005). In bbs-7 and bbs-8 mutants, kinesin-II and OSM-3 separate in the middle segment, which results in IFT-A moving with kinesin-II at a slower rate of 0.5 µm/s and IFT-B moving with OSM-3 at a faster rate of 1.3 µm/s (Ou et al., 2005).

In arl-13 mutants, anterograde IFT movement along distal singlets and retrograde IFT movement occurred at characteristic wild-type rates (Table S1). Because IFT components accumulated abnormally in middle segments, we performed detailed analyses of IFT movement in these regions. The recorded IFT velocities in phasmid cilia were distributed in a normal curve that resembled the previous observation made in amphid cilia (Fig. 2 E; Snow et al., 2004). In arl-13 mutants, the velocity curves of IFT-A and the IFT-A–associated motor kinesin-II shifted to the left (slower) side, whereas those of IFT-B and the IFT-B–associated motor OSM-3 shifted to the right (faster) side (Fig. 2 E).

To determine the molecular basis of the abnormal IFT movement in arl-13 mutants, we subtracted the transport event ratio of wild-type animals from that of arl-13 mutants at each velocity step. Analyses of the distribution curves revealed that in arl-13 cilia, IFT-A and kinesin-II moved at a rate of ~0.5 µm/s, whereas IFT-B and OSM-3 moved at ~1.3 µm/s (see Fig. 5). For all IFT reporters used in our analyses, fewer IFT particles were found to move at the normal rate of ~0.7 µm/s in arl-13 mutants than in wild-type animals. These observations suggest that in ARL-13-deficient cilia, the IFT complex tends to dissociate, and the separated subcomplexes are transported by either kinesin-II or OSM-3 alone (see Fig. 7). The dissociation of IFT-A and IFT-B may account for the ciliogenesis defects and accumulation of IFT-B subcomplexes in arl-13 mutants.

Depletion of ARL-3 partially rescues deformations in ARL-13-deficient cilia
Incomplete dissociation of the IFT complex in arl-13 cilia indicates the involvement of redundant players in this process. arl-3 and arl-6, the C. elegans homologues of human Arl3 and Arl6/BBS3, are promising candidates because they are expressed exclusively in ciliated cells. Similar to ARL-3, ARL-3::GFP does not show detectable movement in cilia (Fig. S2 A), axons, or dendrites (not depicted). In contrast, ARl-6 exhibits typical IFT motility in cilia (Fan et al., 2004). Although ARL-3 was reported to be expressed exclusively in ciliated neurons, with little ciliary localization (Blacque et al., 2005), analysis of our GFP-tagged ARL-3, whose expression was driven by its own 2.5-kb promoter, revealed that a significant amount of ARL-3::GFP localized to cilia (both middle and distal segments; Fig. 3 B). We further analyzed arl-3(m1703) mutants, which carry a genomic deletion that encompasses nearly all of the arl-3 gene (Fig. 3 A). To our surprise, not only were arl-3(m1703) animals superficially wild type, they also had no detectable sign of cilia malformation by the dye-filling assay and IFT markers (Fig. 3, C and D).

To address the possibility that arl-3 negatively regulates ciliogenesis, we generated a transgenic strain that overexpresses ARL-3G72L, a constitutive GTP-binding (dominant active [DA]) mutant. In accordance with our prediction, DA ARL-3G72L overexpression caused Dyf in wild-type phasmid cilia (Fig. 3 C). We characterized the genetic interaction between arl-3 and arl-13. As shown in Fig. 3 C, arl-3; arl-13 double mutants showed less phasmid Dyf, suggesting that depletion of ARL-3 can partly rescue cilia deformations in ARL-13-deficient cilia. This result confirms that ARL-3 negatively regulates ciliogenesis. Consistently, overexpression of DA ARL-3 in arl-13 had an additive effect in causing much more severe Dyf (Fig. 3 C). Further analysis indicated that the abnormal bulblike accumulations of IFT-B components, BBS proteins, and the OSM-3 motor in arl-13 mutants were significantly rescued by the introduction of the arl-3–null allele (Fig. 3 E).

In middle segments of wild-type animals, IFT-A and IFT-B subcomplexes are transported together by kinesin-II and OSM-3 at the intermediate rate of 0.7 µm/s. According to our measurements, the length of the phasmid cilia along which IFT particles move at 0.7 µm/s is ~2.2 µm (Fig. 3 F). In arl-13 mutants, this length decreases to ~1.5 µm. As expected, the length of middle segment can be partly rescued to ~2.0 µm in arl-3; arl-13 double mutants (Fig. 3 F).

Ultrastructural defects in cilia of arl-13 and arl-3; arl-13 mutants
To investigate cilia deformations at the ultrastructural level, we used TEM to examine amphid cilia in various mutants (Fig. 4). Wild-type amphids contained 10 cilia bundled together (Fig. 4 A), whereas arl-13 mutants had only six to eight distal segments, likely as a result of premature truncation of the distal segments of some cilia (n = 8; Fig. 4 B). The remaining distal segments contained normal microtubule singlets. Some truncations in the distal segments continued further, usually resulting in one to two missing middle segments in the worms we sectioned.
an increased number of outer doublets (14 rather than 9 doublets; Fig. 4 S) in a cilium, with most showing B-tubule seam breaks, and some doublets were seen in the ciliary lumen. This structure represents a fused cilium with misplaced doublets in the middle of the ciliary lumen. The *C. elegans* TZ typically contains a circular array of doublet microtubules with y links between the axoneme and ciliary membrane (Fig. 4, M–P, white arrows; Perkins et al., 1986). The number and morphology of amphid cilia in the TZ area of *arl-13* mutants were comparable with those of wild-type animals (Fig. 4, I, J, M, and N).

Several other cilia deformations were observed in *arl-13* mutants. For example, an abnormal membrane distortion (seen in two of eight amphids of *arl-13* mutants; Fig. 4, F and T, black arrows) in middle segments may correlate with the bulblike structure observed in Fig. 2 C. Occasionally, we observed

(Fig. 4 F). In contrast to the normal pattern in middle segments of nine peripheral doublets adjacent to the membrane, misplaced doublets were frequently seen in the ciliary lumen of *arl-13* mutants (Fig. 4, F and T, arrowheads), a phenomenon indicative of a disrupted axoneme architecture or microtubule:membrane connection.

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Consistent with our finding that ARL-3 depletion can rescue various ciliary deformations in *arl-13* mutants, the distal segments of *arl-3; arl-13* mutants were restored to 9–10 cilia, comparable with those of wild-type worms (Fig. 4 C). The abnormal membrane distortion (Fig. 4, F and R, arrows) and uncommon ciliary fusion with broken B-tubules were not
ARL-3 depletion restores IFT-A–IFT-B binding in arl-13 mutants

To gain a mechanistic insight into how ARL-3 depletion rescues ciliogenesis defects in arl-13 mutants, we performed detailed IFT analysis on arl-3; arl-13 double mutants. ARL-3 depletion does not rescue arl-13 ciliogenesis defects by restoring the integrity of the entire IFT particle. Rather, the absence of ARL-3 destabilizes the association between IFT-B and the OSM-3 motor, which indirectly restores binding between IFT-A and IFT-B subcomplexes.

**Figure 5.** ARL-13 and ARL-3 influence the integrity of the IFT macromolecular complex. The IFT transport event ratios of GFP-tagged IFT proteins of wild-type (WT) animals were subtracted from those of arl-3(tm1703), arl-13(gk513), and arl-3; arl-13 mutant animals. Positive value, negative value, and zero (y axis) represent the ratio of more, fewer, or an equal number of particles moving at a given specific velocity step (x axis) in mutants compared with wild-type animals. The simple spline curves in each figure were generated using SigmaPlot software (version 8; Systat Software, Inc.). For all mutants, the curves fit into a bimodal Gaussian distribution, with fewer particles moving at the normal rate of ~0.7 µm/s (transport by kinesin-II and OSM-3 together) and more moving at either ~0.5 (kinesin-II alone) or ~1.3 µm/s (OSM-3 alone), depending on the IFT markers used in the analysis. IFT-A and IFT-B tended to dissociate and move with their own motors in arl-13 mutants, whereas IFT-B tended to dissociate from OSM-3 and move together with IFT-A–kinesin-II in arl-3 and arl-3; arl-13 double mutants. Error bars indicate SEM.
The C. elegans HDAC6 homologue F41H10.6 (hereafter termed HDAC6). The C. elegans hdac6 gene has one available mutant allele, ok3203, which removes ~400 bp of coding sequence and may result in a prematurely terminated protein or alternatively spliced products. hdac6 mutants are superficially wild type and possess normal cilia on the basis of dye-filling experiments. hdac6 gene expression was detected in ciliated amphids and phasmids, nerve cords, intestine, and pharynx (Fig. S3). In ciliated neurons, the GFP::HDAC6 signal is diffuse in the cell body, axon, dendrite, and cilium. Interestingly, GFP::HDAC6 is retained predominantly in the middle segment, like ARL13::GFP (Fig. 6A).

We then generated arl-3; hdac6 double-, arl-3; hdac6 double-, and arl-3; arl-13; hdac6 triple-mutant animals. arl-3; hdac6 double mutants were normal in the dye-filling assay. As shown in Fig. 6B, the hdac6 allele can rescue the ciliogenesis defect in arl-13 to a similar extent as the arl-3 allele. We next asked whether hdac6 and arl-3 act in the same genetic pathway to correct the ciliogenesis defects in arl-13 mutants. Epistasis analysis is commonly used to assess genetic interactions among several genes. We reasoned that if arl-3 and hdac6 act independently or redundantly, we would observe an additive rescue effect in arl-3; hdac6; arl-13 triple mutants. A dye-filling assay showed that the rescue effect in arl-3; hdac6; arl-13 triple mutants is comparable with that in arl-3; arl-13 or hdac6; arl-13 double mutants, suggesting that arl-3 and HDAC6 physically interacts with a BBSome subunit, BBIP10, and is involved in BBIP10-dependent cytoplasmic tubulin acetylation (Loktev et al., 2008). To investigate whether ARL-3 and HDAC6 function in the same pathway to regulate ciliogenesis, we characterized the C. elegans HDAC6 homologue F41H10.6 (hereafter termed HDAC6). The C. elegans hdac6 gene has one available mutant allele, ok3203, which removes ~400 bp of coding sequence and may result in a prematurely terminated protein or alternatively spliced products. hdac6 mutants are superficially wild type and possess normal cilia on the basis of dye-filling experiments. hdac6 gene expression was detected in ciliated amphids and phasmids, nerve cords, intestine, and pharynx (Fig. S3). In ciliated neurons, the GFP::HDAC6 signal is diffuse in the cell body, axon, dendrite, and cilium. Interestingly, GFP::HDAC6 is retained predominantly in the middle segment, like ARL13::GFP (Fig. 6A).

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IFT-A cause Dyf (Fig. 6 B). Therefore, our data reveal that the previously proposed roles for the BBSome and DYF-13 act in the same genetic pathway to regulate cilia formation. Furthermore, TEM analyses confirmed that the distal amphid cilia number can be restored to the wild-type level in 

**Figure 7. Schematic model of ARL involvement in the regulation of IFT stability.** (A) In wild-type [wt] cilia, ARL-13 and ARL-3 stabilize the IFT complex. The holding force between IFT-A and IFT-B is counterbalanced by the dragging force generated by fast-moving OSM-3 motors and slow-moving kinesin-II motors. OSM-3 and kinesin-II coordinate the movement of IFT particles at an intermediate rate. (B) Because of the attenuated association between IFT-A and IFT-B in arl-13 mutants, the holding force is smaller than the dragging force. Some IFT-A and IFT-B are dissociated and moved by their own motors at slow or fast rates. (C and D) In both arl-3 single (C) and arl-3; arl-13 double (D) mutants, the weakened connection between OSM-3 and the IFT-B subcomplex reduces the dragging force that OSM-3 exerts on the rest of the IFT complex, and thereby indirectly restores the binding efficiency of IFT-A and IFT-B.


**Discussion**

Considerable interest exists in further defining the molecular pathways that lead to ciliopathies. Exploring the in vivo molecular mechanisms of conserved disease genes in a simple *C. elegans* model system has helped advance our understanding of the pathophysiology of many ciliopathies, such as autosomal dominant polycystic kidney disease, Merkel–Gruber syndrome, nephronophthisis, and BBS syndromes (Haycraft et al., 2001; Fan et al., 2004; Barr, 2005; Badano et al., 2006; Jauregui et al., 2008; Williams et al., 2008; Bialas et al., 2009).

The cilia-related phenotypes of arl-3 and arl-13 mutants reported in this study are quite reminiscent of those in other organisms. For example, in *L. donovani*, a DA form of ARL-3 (LdARL-3A) caused a truncated flagellum, whereas the dominant-negative form did not affect flagellum biogenesis. *Arl5<sup>−/−</sup>* mice show apparently normal ciliary structure but abnormal ciliary protein transport in photoreceptors. Zebrafish *Arl13b/scorpion* mutants show abnormal cilia formation (Sun et al., 2004), and *Arl13b<sup>−/−</sup>*/<sup>−/−</sup> mice possess shortened cilia with a specific B-tubule closing defect that is similar to what we observed in arl-13 worms (Fig. 4 S; Caspary et al., 2007).

**Effectors of ARL-3 and ARL-13**

Although IFT integrity in arl-3; arl-13 mutants was restored to a similar state as that in arl-3 mutants (Fig. 5; and Fig. 7, C and D), arl-3; arl-13 mutants still display mild ciliogenesis defects. One hypothesis to explain this observation is that the effectors of ARL-13 may have two distinct functions: one related to regulating IFT integrity and the other a non-IFT-related function, such as membrane–microtubule architecture. This hypothesis is supported by the observation with TEM that arl-3; arl-13 mutants possess misplaced doublets. ARl-3 depletion indirectly restores IFT but is not able to correct other non-IFT-related ciliogenesis defects; therefore, we did not see a complete rescue effect for the arl-3; arl-13 double mutants.

The cellular effectors of ARL-3 and ARL-13 are poorly understood. However, experiments on ARL mutants with altered GTP-binding capacity indicate that normal ciliogenesis requires the GTPase activities of both ARL-3 and ARL-13 (Fig. 2 A and Fig. 3 C). Identification of the putative GTPase exchange factor and GTPase-activating protein that bind to ARL-3 and ARL-13 will be critical to fully dissect their roles in cilia biogenesis and IFT regulation. Remarkably, the proposed roles of ARL-13 and ARL-3 in IFT regulation resemble the roles proposed for the BBSome and DYF-1 in this system (Ou et al., 2005). It would be informative to study how ARLs, BBS
proteins, and DYF-1 converge in the same pathway to regulate ciliogenesis and IFT.

The C-terminal PRD of ARL-13 is indispensable for its specific targeting to the doublet segment of the cilium. The PRD and its major interacting domain, SH3 (Src homology 3), have been shown to be involved in various protein–protein interactions. Interestingly, a PRD is found in many microtubule-associated proteins such as MAP2, τ, and MAP4 and is required for the activities of microtubule-associated proteins in microtubule binding and assembly (West et al., 1991; Kanai et al., 1992; Ferralli et al., 1994; Tokuraku et al., 1999). Additionally, the PRD of the GTPase dynamin regulates its effector binding and microtubule association (Gout et al., 1993; Hamao et al., 2009).

In our experiments, overexpression of the ARL-13 PRD domain induced a dominant-negative effect on ciliogenesis (Fig. 2 A). Overexpression of the ARL-13 NT truncation, which lacks the PRD domain but contains the entire GTPase domain, could also cause Dyf in the phasmid cilia of wild-type animals. Similarly, the arl-13(tm2322) allele is an in-frame deletion and encodes a truncated ARL-13 protein that lacks the PRD domain but retains the intact GTPase domain (Fig. 1 A). Both arl-13(gk513) and arl-13(tm2322) animals show comparable ciliogenesis defects (unpublished data). Remarkably, contrary to our observation that none of the IFT reporters interfere with ciliogenesis in arl-13 (gk513) mutants, IFT transgene overexpression can enhance the ciliogenesis defect of arl-13(tm2322) mutant cilia (Cevik et al., 2010). These findings provide support for the view that the PRD domain of ARL-13 is critical for both ciliary targeting and cilia formation, and loss of the PRD domain could result in a truncated ARL-13 protein with deregulated GTPase activity. Thus, characterization of the ciliary proteins that bind the ARL-13 PRD domain is of great interest.

What are HDAC6’s substrates in ciliogenesis?

We demonstrate in this study that ARL-3 acts through the deacetylase HDAC6 to regulate cilia formation in C. elegans. HDAC6 deacetylates not only histones (Grozinger et al., 1999) but also α-tubulin (Hubbert et al., 2002), Hsp90 (Kovacs et al., 2005), and cortactin (Zhang et al., 2007). Inhibition of HDAC6 enzymatic activity causes increased transport of the kinesin-1 cargo complex along microtubules in neuronal cells (Reed et al., 2006). Reminiscent of our findings, activation of HDAC6 can promote cilia disassembly, and loss of HDAC6 activity selectively stabilizes cilia in human retinal epithelial cells (Pugacheva et al., 2007).

Currently, HDAC6 is thought to regulate microtubule-dependent processes by deacetylating α-tubulin. However, this model is inaccurate for the C. elegans ciliogenesis pathway because mec-12, the only α-tubulin capable of being acetylated, is not expressed in ciliated amphids and phasmids (Fukushige et al., 1999). Another HDAC6 substrate, Hsp90, was shown to associate with β-tubulin and may be involved in stabilizing tubulin polymerization in cilia (Takaki et al., 2007). However, worms with a null allele of daf-21, the C. elegans homologue of Hsp90, possess normal cilia in our analysis (unpublished data). These data rule out α-tubulin and Hsp90 as candidate HDAC6 substrates during ciliogenesis in C. elegans. On the basis of our model, we propose that HDAC6 deacetylates an unknown adaptor protein in IFT-B–OSM-3 subcomplexes.

Non-IFT-related roles for ARL-3 in cilia formation

ARL-3 appears to have non-IFT–related functions in cilia. Overexpression of DA ARL-3 in arl-13 mutants produces a new defect in that some cilia are missing in the TZ area (Fig. 4 L). This striking phenotype is not the result of compromised IFT in middle segments because previous TEM analyses showed that TZs can form normally in all IFT mutants examined (Perkins et al., 1986). Because C. elegans and vertebrate ARL-3s are cytosolic (Fig. 3 B; Behnia et al., 2004; Setty et al., 2004) and at least one IFT protein (IFT20) moves between the Golgi apparatus and adjacent basal body (Follit et al., 2006), the overexpression of DA ARL-3 could interfere with normal protein transport between the Golgi and TZ. When this interference is combined with the already compromised ciliogenesis of arl-13 mutants, the normal TZ formation could be disrupted. A possible ARL-3 effector in this pathway is the retinitis pigmentosa 2 (RP2) protein. In the mammalian retina, RP2 interacts with GTP-bound Arl3 and functions as a GTPase-activating protein (Grayson et al., 2002; Veltel et al., 2008). K08D12.2, the C. elegans homologue of RP2, is expressed exclusively in ciliated cells and localizes specifically around TZs but not along the axoneme (Blacque et al., 2005). It would be informative to examine whether ARL-3 and RP2 form a functional complex and play a role in the early stage of ciliogenesis.

In conclusion, our findings reveal a novel coordination mechanism for ARL-13 and ARL-3 in cilia formation. Considering the strikingly similar mutant phenotypes across different species, the ciliary roles of ARL-13 and ARL-3 identified in our study are probably evolutionarily conserved in other organisms. The next challenge is to elucidate the effectors of ARL-13 and ARL-3 and the substrates of HDAC6 involved in regulating ciliogenesis.

Materials and methods

C. elegans mutant alleles and strains

Nematodes were raised using standard conditions described previously (Brenner, 1974). N2 worms represented the wild-type animals in all assays. All strains used in this paper are listed in Table S2.

Dye-filling assay

The stock Dil [2 mg/ml in dimethyl formamide; Invitrogen] was diluted 1:200 in M9. Worms were incubated in diluted dye for 1 h at room temperature. After incubation, the animals were washed at least three times with M9 and observed using a fluorescence microscope (M2Bio; Carl Zeiss, Inc.).

Microscopy

Animals were raised at 20°C and imaged using standard C. elegans slide mounts and a Plan Apochromat 60× 1.49 NA oil objective [Nikon] on an imaging microscope (TE 2000-U; Nikon).

IFT measurement

We performed all IFT analyses in phasmids for easier observation. IFT motility was observed using a Plan Apochromat 100× 1.49 NA oil total internal reflection fluorescence objective [Nikon]. In wild-type worms, the IFT velocities in phasmid cilia were identical to those reported in amphid cilia (Table S1). Motility stacks were recorded using a charge-coupled device camera (QuantEM 512SC; Roper Industries), and kymographs were produced.
using MetaMorph software (MDS Analytical Technologies). Worms were anesthetized in a drop of M9 containing 10 mM levamisole, transferred to an agarose mount slide, and imaged immediately.

TEM
For TEM, worms were fixed in 2.5% glutaraldehyde in cacodylate buffer on ice. Heads were cut off, moved to fresh fixative, and held overnight at 4°C. Animals were rinsed in buffer and stained with 1% osmium tetroxide in cacodylate buffer for 1 h at 4°C. After embedding small groups of worms in agarose, the specimens were dehydrated and embedded in Embed®12 resin according to the general procedures described previously [Hall, 1995]. Thin sections were collected on a diamond knife and poststained before viewing on an electron microscope (JE&M-1400, JEOL).

Online supplemental material
Fig. S1 shows that arl-3 and arl-13 are expressed exclusively in ciliated cells. Fig. S2 shows that LRAL-3 and LRAL-13 show no IFT motility. Fig. S3 shows that hda-oc is expressed in ciliated cells. Fig. S4 shows that LRAL-3 localizes normally in arl-13 mutants, and LRAL polymorphism appears normal in ARAL-3Q72L[ΔX12] and arl-13 animals. Table S1 shows IFT velocities of various IFT markers in arl-3 and arl-13 single mutants and arl-3 arl-13 double mutants. Table S2 shows all of the strains used in this study. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.20091201.1/D1C.

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