Functional Genomics of the Cilium, a Sensory Organelle

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Summary

Cilia and flagella play important roles in many physiological processes, including cell and fluid movement, sensory perception, and development [1]. The biogenesis and maintenance of cilia depend on intraflagellar transport (IFT), a motility process that operates bidirectionally along the ciliary axoneme [1, 2]. Disruption in IFT and cilia function causes several human disorders, including polycystic kidneys, retinal dystrophy, neurosensory impairment, and Bardet-Biedl syndrome (BBS) [3-5]. To uncover new ciliary components, including IFT proteins, we compared C. elegans ciliated neuronal and nonciliated cells through serial analysis of gene expression (SAGE) and screened for genes potentially regulated by the ciliogenic transcription factor, DAF-19 [6]. Using these complementary approaches, we identified numerous candidate ciliary genes and confirmed the ciliated-cell-specific expression of 14 novel genes. One of these, C27H5.7a, encodes a ciliary protein that undergoes IFT. As with other IFT proteins, its ciliary localization and transport is disrupted by mutations in IFT and bbs genes. Furthermore, we demonstrate that the ciliary structural defect of C. elegans dyf-13(mn396) mutants is caused by a mutation in C27H5.7a. Together, our findings help define a ciliary transcriptome and suggest that DYF-13, an evolutionarily conserved protein, is a novel core IFT component required for cilia function.

Results and Discussion

Identification of Candidate Ciliary Genes through SAGE and a Bioinformatic Screen for Genes Potentially Regulated by X Box Promoter Elements

In the first approach, we used LongSAGE to obtain a C. elegans ciliated-cell transcriptome. Ciliated neurons expressing GFP from the ciliated-cell-specific bbs-1 gene promoter [5] were isolated and used to construct a LongSAGE library as described [7, 8]. To identify candidate ciliary genes, we compared our ciliated-cell SAGE transcriptome to pan-neural-, muscle-, and gutspecific transcriptomes, all normalized to 50,000 tags (available at http://www.bcgsc.ca). Because ciliated neurons account for only ~20% of all embryonic neurons, genes expressed predominantly in ciliated neurons should be represented by more SAGE tags in the ciliated-cell (C) transcriptome than in the pan-neural (P) transcriptome. Similarly, ciliated-cell-specific transcripts should be underrepresented in the nonciliated muscle (M) and gut (G) transcriptomes. Table S1 (in the Supplemental Data available with this article online) presents the SAGE data, sorted by genes with the most favorable ciliated-cell SAGE tag enrichment factors across all three comparison groups (C/P, C/M, and C/G). As expected, the top-ranked genes from Table S1 contain numerous cilia-related genes, including IFT-related genes (daf-19, bbs-1, che-2, che-13, and osm-6), structural components (tba-9, tba-6, and mec-12) and motor components (F41G4.1, axonemal dynein; dli-1, cytoplasmic dynein; and klp-11, kinesin). To assess the specificity of our SAGE approach in detecting ciliatedcell-specific genes, we screened Table S1 for genes known to be expressed predominantly in ciliated cells; 25 of 42 such genes found (60%) were enriched for SAGE tags in the ciliated-cell transcriptome versus the pan-neural, muscle, and gut transcriptomes (Table S2). These data suggest that Table S1 likely contains many excellent candidate ciliary genes. To estimate the proportion of false positives in Table S1, we investigated the top 200 genes and found 51 genes with known expression data; 34 of these genes (67%) are expressed in nonciliated cells, indicating a relatively high but acceptable false-positive rate. On the basis of these findings, we sought to further refine our search for candidate cilia-related genes with a complementary approach.

We performed a genome-wide bioinformatic screen for genes potentially regulated by X boxes. In C. elegans, these conserved 14 bp regulatory elements (usually 50-150 bp upstream of start codons) are bound by DAF-19, the RFX-type transcription factor that regulates ciliary gene (IFT and bbs) expression [5, 6, 9, 10]. We derived an improved X box consensus sequence for our screen by confirming the ciliated-cell-specific expression of several candidate X-box-containing genes identified in a previous, more preliminary study [9]. Using transcriptional-GFP reporters, we observed ciliated-cellspecific expression for five novel genes, namely K03E6.4, C48B6.8, C47E8.6, ZK328.7a, and F19H8.3 (Figure S1; data not shown for F19H8.3). Figure S1 also shows the ciliated-cell-specific pattern for Y110A7A.20 and K08D12.2, which were not shown in our previous study [9]. Importantly, the expression of ZK328.7a, Y110A7A.20, and C48B6.8 was found to be dependent on DAF-19 (Figure S2). Using an updated Hidden Markov Model X box profile based on 22 sequences, including the novel X-box-containing genes discussed above (Table S3), we scanned the C. elegans genome for X boxes within 1500 bp upstream of start codons. A total of 1572 putative X-box-containing genes with human homologs were identified (Table S4). As expected, the greatest number of high-scoring (>16.0) X boxes occurred between -60 and -120 bp (Figure S3A), which coincides with the canonical position [6].

As described above for the SAGE data, we found that the number of false positives in Table S4 is relatively high, but the dataset still contains a large proportion of bona fide ciliary genes. Of the top 200 ranked candidate X box genes with known anatomic expression data (64 genes), 27 (or 42%) are expressed in ciliated cells.

To identify our best candidate ciliary genes, we considered genes with enrichment of SAGE tags in the ciliated SAGE dataset versus the pan-neural, muscle, and intestine datasets (Table S1), and with putative X boxes within 250 bp of start codons (Table S4). Using these criteria, we present 46 such genes in Table 1. Also included are seven additional genes (at the bottom of Table 1), which rank highly in either the SAGE or X box datasets. As expected, Table 1 contains many known IFT-related genes, including che-2, che-13, osm-1, osm-5, osm-6, F32A6.2 (IFT81), bbs genes, and daf-19. Other excellent candidate and known ciliary genes include K08D12.2 (homolog of retinitis pigmentosa 2), which is expressed predominantly in ciliated neurons (Figure S1), and Y32G9A.6, an ankyrin-repeat-containing protein that is homologous to inversin, which is linked to cilia-associated situs inversus and cystic-kidney phenotypes [11]. Table 1 also includes uncharacterized genes identified as candidate ciliary genes in other genomic and proteomic screens [10, 12, 13]; these genes include C27H5.7a (TPR containing), C54G7.4 (WD repeat containing), K07G5.3 (Ca2+ binding domain containing), and two B9-protein-domaincontaining genes (Y38F2AL.2 and K03E6.4), as well as several genes not previously associated with a ciliary function. C27H5.7a is particularly interesting because it was also recently found by Colosimi et al. [14] to be expressed in the AWB ciliated olfactory neuron. Importantly, the number of false-positive genes in Table 1 is relatively low; 13 of the 20 genes from Table 1 with known anatomic expression data (WormBase [http://www.wormbase.org] and Figure S1) are expressed in ciliated cells. Along with GFP-expression data presented below, we estimate the false-positive rate for the top 46 genes in Table 1 to be approximately 26% (7 of 27 genes), which is significantly lower than that found for the individual SAGE (67%) and X box (58%) analyses.

Expression Analysis of Candidate Ciliated-Cell-Specific Genes

Nine new candidate ciliated-cell-specific genes were chosen for expression analysis on the basis of strong SAGE and X box data (Table 1; seven genes) or a high ranking in the individual X box table (Table S4; two genes). Using transgenic worms bearing promoter-GFP reporters, we found that all nine genes demonstrated a ciliated-cell-specific expression pattern (Figure 1; data not shown for C02H7.1). Transcriptional gfp reporters for C27H5.7a (unknown function), Y38F2AL.2 (B9 domain containing), H01G02.2 (serine/threonine kinase), C54G7.4 (WD domain containing), D1009.5 (dynein light chain), and Y32G9A.6 (inversin-like) produced strong GFP signals in most ciliated cells, including amphid, phasmid, and labial-quadrant neurons. Similarly, F54C1.5a (TPR domain containing) and K07G5.3 (Ca2+ binding domain containing) expressed in most amphid, both phasmid, and several labial-quadrant neurons. That the expression of the gfp reporters was not detected in nonciliated cells suggests that these novel genes have ciliary functions. It is notable that all of the genes selected from Table 1 for expression analysis demonstrated a ciliated-cell expression pattern. This is a very strong indication that Table 1 is highly enriched for novel cilia-related genes.

Cellular Localization of Candidate Ciliary Proteins

To investigate the possible functions of some of the ciliated-cell-specific proteins identified above, namely C27H5.7a, Y37E3.5, K08D12.2, and F19H8.3, we examined their cellular localization with translational gfp reporters (Figure S4A). Interestingly, GFP-tagged C27H5.7a (unknown function) and Y37E3.5 (GTP binding protein, ARL2-like-1) associate with ciliary axonemes in both head (amphids, labials) and tail (phasmids) neurons, with relatively little localization to cell bodies, axons, or dendrites. In addition, C27H5.7a::GFP localizes to the transition zones (basal bodies at the base of cilia). GFPtagged Retinitis Pigmentosa 2 homolog K02D12.2 appears to envelop the transition zones in a ring-like fashion (Figure S4A), indicating a possible role at the base of cilia. GFP-tagged ARL-3 (F19H8.3) is found throughout the cell, with little ciliary localization even though its expression is limited to ciliated cells.

C27H5.7a Is a Novel IFT-Associated Protein

A major focus of our functional genomic studies is to uncover novel core ciliary proteins, including IFT com-

Table 1. SAGE and X Box Data Identify a Set of Strong Candidate Cilia-Related Genes

Gene	SAGE Data				Xbox Data			Other Studies				Annotation /	
Model	Name	С	Р	М	G	Sequence	Pos.	Score	Α	В	С	D	Description
Y105E8A.5	bbs-1	19	1	0	0	GTTCCCATAGCAAC	98	19.29	/	√			BBS1
F59C6.7	che-13	26	3	1	1	GTTGCTATAGCAAC	73	17.70	/	✓	/		IFT57
C27H5.7a		8	0	0	1	GTCTCCATAGCAAC	102	19.07	/	✓	/	/	Uncharacterized
T06G6.3		7	1	0	0	GTTGCCATGGCAAA	59	16.11					Myosin heavy chain-
		_		_					,				like
Y38F2AL.2	oom 6	6 6	1 1	0	0	GTTGCCGTGGCAAC	65 99	16.11 18.68	✓ ✓	✓ ✓			B9 domain containing IFT52
R31.3 F32A6.2	osm-6	5	1	0	0	GTTACCATAGTAAC GTTGCCCTGGTAAC	99 71	16.85	✓ ✓	~	✓		IFT81
F38G1.1	che-2	5	1	1	1	GTTGTCATGGTGAC	129	16.30	V	_	V		IFT80
R13G10.2	amx-1	5	1	0	1	GCTACCATGACAAC	144	14.52	•	v	•		Monoamine oxidase
F35D2.4		5	1	0	0	ATCACCATGACAAC	158	12.85					Uncharacterized
M04G12.3	gcy-34	4	0	1	0	GTCTTCACAGCAAC	122	11.09		✓			Guanylate cyclase
K10G6.4		4	1	0	0	GTCACCATGCCAAT	139	12.85					Uncharacterized
Y41G9A.1	osm-5	8	2	1	0	GTTACTATGGCAAC	114	19.79	✓	✓	✓		IFT88
W02B12.3c	rsp-1	7	0	2	1	GTTGTCATGGCTAC	131	15.18					Splicing factor
F33H1.1b	daf-19	28	8	6	7	GTTTCCATGGAAAC	108	19.61					RFX transcription
			_	_	_								factor
F08C6.4	sto-1	30	9	8	6	TTTTCCTTGGTAAC	201	12.69	,				Stomatin
K03E6.4	l d	3	1	0	0	GTTCCCTTGGCAAC	83	17.11	/				B9 domain containing
F13B9.5	ksr-1	3	1	1	0	TTTTTCTTGGCAAC	104	10.65					Kinase regulator of Ras
Y32G9A.6		3	1	0	0	GTTGTCAGGGTAAC	186	13.52					Inversin-like
Y41D4A.4		6	1	0	2	GTTGCCACGTTAAC	140	12.46					EMP70
F32A6.5	sto-2	6	1	2	1	GTTTCCTTAAAAAC	181	11.48					Stomatin
F28E10.1c		11	4	0	2	GTCGCCAGAGCAAC	80	13.09					Titin/Connectin
F20D12.3	bbs-2	8	3	0	1	GTATCCATGGCAAC	93	20.02	✓				BBS2
ZC84.6		5	2	1	0	GTTTATTTGGTAAC	99	11.11					Serine protease
V074440		•	•	•	•	0.000.000.000.000	470	40.00					inhibitor
Y37A1A.3		2 2	0 1	0	0 1	GTCAATATGACAAC	178	10.26					Glucose transporter
B0228.5a	+ 0	2	1	0	0	GTTACCATATCAGC	16	11.18		,			Thioredoxin
B0432.5a	cat-2	2	1	1	0	GTCCCCACGCCAAC	36	10.97		√,			Tyrosine hydroxylase
F56H11.1c K08D12.2	fbl-1	2	1	0	0	GTTTCCATACCAAC	76 83	10.09 20.74	/	✓			Fibulin Retinitis pigmentosa 2
C27F2.1		2	1	1	0	GTTTCCATAGCAAC GTCACCATAGCAAC	83	18.41	V				Uncharacterized
T27B1.1	osm-1	2	0	0	1	GCTACCATGGCAAC	85	17.91	/	_	/		IFT172
C09E10.2b	dgk-1	2	1	0	0	GTACTCATGCTAAC	131	10.30	~	V	~		Diacylglycerol kinase
H01G02.2	ugk-1	2	1	1	0	GTCTCCATGCTAAC	159	16.96					Serine/Threonine
1101002.2		2	•	'	U	atotooataaoaao	100	10.30					kinase
C26B9.5		2	0	0	1	GTTTGCATAGTCAC	174	10.41					Serine protease
T01D1.6	abu-11	2	0	0	0	GTCAACATAACAAC	229	10.56					Keratin associated
K07G5.3		6	3	1	1	GTTGCCATAGCGAC	70	17.30		✓			Uncharacterized
K06H7.3		6	3	1	1	TTTTCCATAATAAC	238	11.48					Ankyrin repeat protein
R166.3		6	3	2	3	ATCTCCTTGGCAAC	248	13.43					AMME Syndrome 1
R11A8.2		7	4	0	0	GTTTCCATATTAAT	225	11.48					G patch and KOW
ZK1151.1c	vab-10	7	4	1	0	GTTTTTATGGCAAA	235	12.52					motifs MACF1
R01H10.6	bbs-5	19	11	2	3	GTCTCCATGGCAAC	64	20.35	/	/			BBS5
T24D1.1b	sqv-5	5	3	0	0	GTTCCTTCGGCAAC	33	11.00	•	v			Chodroitin synthase 2
F59D12.1	041 0	3	2	1	0	ATTACCATAGTTAC	170	12.92					Transmembrane
			_	•		7.1. 11.10 07.111.10							protein
C02H7.1		3	2	0	0	GTCTCCATGACAAC	194	16.96	\checkmark	\checkmark			Uncharacterized
D1009.5		6	4	2	2	GTTGCCATGACAAC	76	17.18	✓	\checkmark			Dynein light chain
C48B6.8		6	4	2	0	GTTTCCATGACAAC	80	18.63					B1 protein
F08B12.1		13	1	1	1				,				Prominin 1
ZK520.1		8	2	1	2	077000470004	470	47.44	/	√ ,			WD-repeat domain 19
C54G7.4		4	4	0	0	GTTGCCATGGCAAT	172	17.11	✓,	✓,			WD-repeat protein 35
F54C1.5a		3	3	1	2	GTTACCATGGATAT	105	15.50	/	√			TPR repeat containing
ZK328.7a		1	1	1	0	GTTACCATGGCAAT	88	17.91	/	,			TPR repeat containing
F19H8.3 C47E8.6		1	6 0	2 0	1 0	GTCTCTATGGTAAC	150	17.35 17.98		✓			ARL3 Uncharacterized
047 LO.0		0	U	U	U	GTTACCATGCCAAC	108	17.90					Officialactefized

Shown are all genes (46 in total) that are enriched for SAGE tags in the ciliated-cell transcriptome (C), with \geq 1.5-fold enrichment versus the pan-neural transcriptome (P) and \geq 2.0-fold enrichment versus the muscle (M) and gut (G) transcriptomes, and with a canonically-positioned (Pos) putative X box (i.e., within 250 bp of the start codon). Only genes with homology to a human gene (Blast E-value < e^{-10}) are represented. Genes are sorted according to the level of SAGE tag enrichment in the ciliated-cell transcriptome versus the other three transcriptomes. The seven candidate cilia-related genes at the bottom of the table are included on the basis of their high ranking in the individual SAGE or X box tables (Tables S1 and S4, respectively). Check marks indicate genes identified as candidate ciliary genes in three previous genomics and proteomics studies, namely (A) Li et al. [10], (B) Avidor-Reiss et al. [12], and (C) Ostrowski et al. [13], or as a sensory-neuron-type-specific gene ([D] Colosimo et al. [14]).

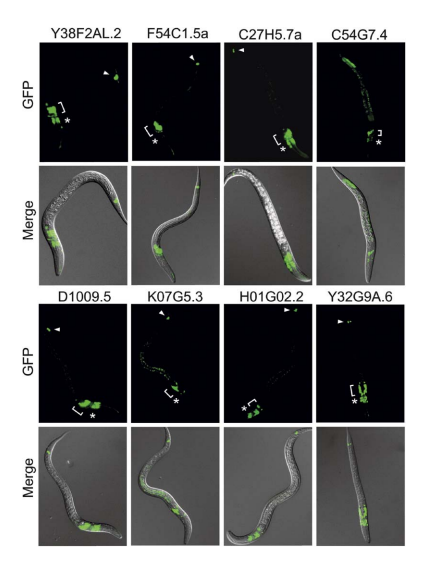


Figure 1. Expression Analysis of Candidate Cilia-Related Genes Identifies New Genes Expressed Exclusively in Ciliated Cells

Shown are whole-worm GFP and GFP/DIC merged images of transgenic animals expressing the indicated transcriptional *gfp* reporter. Expression is restricted to ciliated cells such as the amphid neurons (bracket), the labial-quadrant neurons (asterisk), and the phasmid-tail neurons (arrowhead). Note that for some markers (e.g., C54G7.4 and K07G5.3), the weak signals along the midbody are due to the nonspecific autofluorescence of gut granules.

ponents. First described in *Chlamydomonas* [15], IFT involves the kinesin- and dynein-dependent bidirectional movement of IFT particles and associated cargo along ciliary axonemes [1–3, 15]. In *C. elegans*, anterograde transport of IFT particles along the middle segment of cilia is driven cooperatively by two Kinesin-2 motors (heterotrimeric Kinesin-2 and homodimeric OSM-3) and by OSM-3 alone along the distal segments [16]. IFT particles have been shown biochemically to be composed of two large protein complexes (A and B) containing at least 16 proteins in total [17, 18].

Using time-lapse microscopy, we examined animals expressing C27H5.7a::gfp and found that the GFP-tagged protein exhibited IFT along ciliary axonemes of head and tail neurons (Figure S4B; Document S2; Movies S1 and S4). Anterograde transport rates along the middle and distal segments were determined to be 0.70 \pm 0.11 $\mu m/s$ (n = 245) and 1.25 \pm 0.15 $\mu m/s$ (n = 255), respectively, with the retrograde transport rate estimated to be 1.32 \pm 0.2 $\mu m/s$ (n = 5). Remarkably, these transport rates for C27H5.7a::GFP are almost identical to those previously reported for the Kinesin-2 motors and IFT particle components [1, 16], indicating

that C27H5.7a is transported with IFT particles by heterotrimeric Kinesin-2 and OSM-3 within the middle segments of ciliary axonemes and by OSM-3 kinesin alone within the distal segments.

We then examined the behavior of C27H5.7a::GFP in two IFT mutant strains, namely osm-5(p813) and che-11(e1810), both of which possess truncated cilia and display abrogated IFT. In these mutants, C27H5.7a:: GFP displayed little or no detectable IFT and accumulated in the dendrite and transition zones of the osm-5 mutant, as well as within the transition zones and along the ciliary axonemes in the che-11 background (Figure 2A; Document S2; Movies S2, S3, S5, and S6). The dependence of C27H5.7a on osm-5 and che-11 for its normal localization and motility strongly suggests that C27H5.7a is part of the motor-IFT particle-cargo complex. To provide further evidence that C27H5.7a is likely a core component (and not cargo) of IFT particles, we studied its behavior in bbs mutant backgrounds. Previously, we observed that GFP-tagged CHE-2 often accumulated near the midpoint and distal segments of ciliary axonemes in bbs mutants [19]. We now show that three other GFP-tagged IFT components, namely

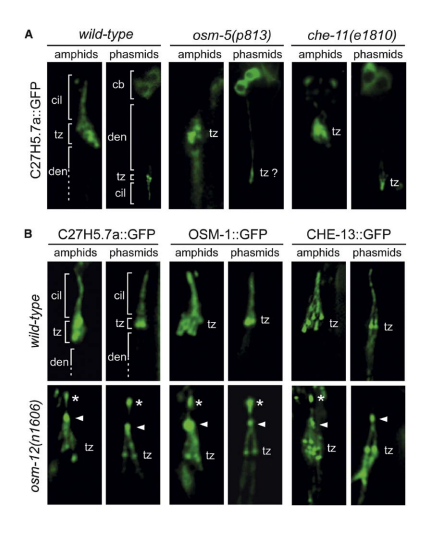


Figure 2. The Ciliary Localization of C27H5.7a Is Disrupted by Mutations in IFT and bbs Genes

Shown are one set of amphid and phasmid cilia, with the ciliary axonemes (cil), transition zones (tz), dendrites (den), and cell bodies (cb) indicated in the left-hand panels. All images are similarly sized and oriented, with the tz positions indicated in all panels.

(A) Localization of GFP-tagged C27H5.7a in wild-type, osm-5(p813), and che-11(e1810) mutant backgrounds. In the wild-type, C27H5.7a::GFP localizes at the transition zones and along ciliary axonemes, whereas in osm-5(p813) and che-11(e1810) mutants, C27H5.7a::GFP accumulates at transition zones in osm-5 mutants and along the stunted ciliary axonemes of che-11 mutants. The question mark indicates that the tz position cannot be accurately identified.

(B) C27H5.7a::GFP mislocalizes along bbs-7 (osm-12(n1606)) mutant cilia, displaying accumulation at the midpoint of the ciliary axonemes (arrowhead) and within the distal segment (asterisk). Similar mislocalization along bbs-7 mutant cilia is also observed for four GFP-tagged IFT proteins, namely OSM-1, CHE-13, OSM-6, and CHE-2 (see Figure S5 for OSM-6::GFP and CHE-2::GFP images).

CHE-13, OSM-1, and OSM-6, demonstrate the same phenotype in a *bbs-7* mutant (*osm-12(n1606)*) (Figure 2B; data for CHE-2 and OSM-6 shown in Figure S5). Interestingly, when expressed in the *bbs-7* or *bbs-8*

mutant backgrounds, C27H5.7a::GFP displays the same mislocalization phenotype as OSM-1, OSM-6, CHE-2, and CHE-13 (Figure 2B; Document S2; Movies S7–S16; data not shown for *bbs-8* mutant), suggesting

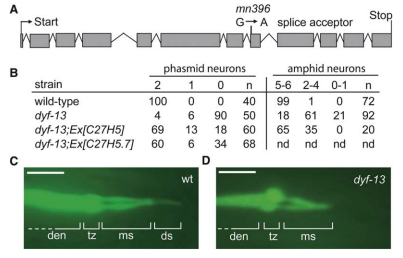


Figure 3. Mutation in C27H5.7a Causes the Structural Cilia Defect in *dyf-13(mn396)* Mutants

- (A) A schematic of the putative exon/intron structure of C27H5.7a shows the guanine-toadenine splice acceptor mutation in dyf-13(mn396).
- (B) The dye-uptake-defective phenotype (Dyf) of dyf-13(mn396) mutants is rescued by the C27H5 cosmid and by a transgene containing the wild-type C27H5.7a gene. Shown are the percentages of sensory organs with normal dye-fill in the indicated number of amphid and phasmid neurons. "n" denotes number of sensory organs scored and "nd" denotes not determined.

(C and D) The cilia of *dyf-13(mn396)* mutants are short, lacking distal portions. Shown are GFP images of animals expressing a *ceh-23* promoter::*gfp* transgene in the phasmid neurons. Anterior is to the left. Note that the

phasmid cilia of dyf-13 mutants are shorter than those of wild-type animals. The following abbreviations were used: den, dendrite; tz, transition zone; ms, middle segment; and ds, distal segment. The scale bar represents 5 μ m.

that C27H5.7a is closely associated with IFT particle components and does not represent cargo.

The dyf-13 Mutant Contains a Mutation in C27H5.7a and Possesses Abnormal Cilia

To provide additional evidence that C27H5.7a is a core ciliary component, we searched existing sensory mutants for mutations in this gene. Using this approach, we found that dyf-13(mn396) animals contained a guanine-to-adenine mutation in the acceptor site of intron 7 of C27H5.7a (Figure 3A). dyf-13 mutants are defective in sensory neuron dye filling and chemosensation [20]. Specifically, only 18% of amphid and 4% of phasmid sensory organs displayed wild-type dye filling (Figure 3B). This phenotype, comparable to that of IFT and bbs mutants [19-21], suggests that ciliary structures are compromised. Using a ceh-23 promoter::gfp reporter (gift of G. Garriga) that illuminates the cell bodies, dendrites, transition zones, and cilia of the phasmid neurons, we found that the cilia of dyf-13 animals were indeed malformed (Figures 3C and 3D). Specifically, the average length of the cilium in dyf-13(mn396) animals was 6.7 \pm 0.8 μ m (n = 20) in comparison to 8.6 \pm 1.6 μ m (n = 20) in wild-type animals, and the neurons appeared to lack the distal portion of the cilia. This cilia defect is similar to that observed in animals harboring mutations in osm-3, an IFT-specific kinesin required to build the distal segment [16, 21]. Importantly, the dyefilling defect of dyf-13(mn396) mutants was similarly rescued in three independent lines expressing either the C27H5 cosmid or a transgene bearing the wild-type C27H5.7 gene (Figure 3B; data shown for one rescued line in each case).

On the whole, our findings provide strong evidence that DYF-13 (C27H5.7a) is an IFT protein that is important for the formation and function of cilia. Consistent with this conclusion, an uncharacterized human homolog (FLJ12571) with 37% amino acid identity and 59% similarity throughout its length exists, and counterparts are found in all ciliated organisms (including *Chlamydomonas reinhardtii*), but not in nonciliated organisms such as *Arabidopsis thaliana* or *Saccharomyces cerevisiae*.

Supplemental Data

Detailed Experimental Procedures and several supplemental figures, movies, and tables are available at http://www.current-biology.com/cgi/content/full/15/10/935/DC1/.

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to *dyf-13(mn396)* (phenotypic characterization and cloning of the mutation) should be addressed to S.S.

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