

Cycling of the signaling protein phospholipase D through cilia requires the BBSome only for the export phase

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The BBSome is a complex of seven proteins, including BBS4, that is cycled through cilia by intraflagellar transport (IFT). Previous work has shown that the membrane-associated signaling protein phospholipase D (PLD) accumulates abnormally in cilia of *Chlamydomonas reinhardtii* *bbs* mutants. Here we show that PLD is a component of wild-type cilia but is enriched ~150-fold in *bbs4* cilia; this accumulation occurs progressively over time and results in altered ciliary lipid composition. When wild-type BBSomes were introduced into *bbs* cells, PLD

was rapidly removed from the mutant cilia, indicating the presence of an efficient BBSome-dependent mechanism for exporting ciliary PLD. This export requires retrograde IFT. Importantly, entry of PLD into cilia is BBSome and IFT independent. Therefore, the BBSome is required only for the export phase of a process that continuously cycles PLD through cilia. Another protein, carbonic anhydrase 6, is initially imported normally into *bbs4* cilia but lost with time, suggesting that its loss is a secondary effect of BBSome deficiency.

Introduction

Bardet-Biedl syndrome (BBS; OMIM accession no. 209900) is a rare inherited disorder characterized by retinal degeneration, anosmia, kidney anomalies, polydactyly, hypogonadism, and obesity (Beales, 2005; Blacque and Leroux, 2006; Zaghoul and Katsanis, 2009; Sheffield, 2010). The phenotype of BBS is indicative of defects in the function of cilia, and specifically in cilia-mediated signaling. Mutations in at least 12 genes (*BBS1*–*BBS12*) cause the syndrome (Jin and Nachury, 2009). Seven of the *BBS* gene products (*BBS1*, 2, 4, 5, 7, 8, and 9) form a biochemically stable complex, the BBSome (Nachury et al., 2007). The BBSome subunits are well conserved in organisms with cilia, indicating that the BBSome fulfills an important ciliary function. Recently, we identified *Chlamydomonas reinhardtii* mutants for *BBS1*, *BBS4*, and *BBS7*. Because large quantities of cilia can be isolated from *C. reinhardtii*, it was possible to carry

out a direct biochemical comparison of wild-type and *bbs* mutant cilia. (Because cilia and flagella are essentially identical organelles, here, we refer to the two flagella of *C. reinhardtii* as cilia.) Remarkably, loss of the BBSome has little effect on the overall composition of cilia or the ciliary axoneme; rather, a small subset of membrane-associated proteins, several of which are predicted to have signaling function, are present at abnormal levels in *bbs* cilia (Lechtreck et al., 2009). A redistribution of ciliary membrane proteins is also characteristic for *Bbs* knockout mice (Berbari et al., 2008b; Domire et al., 2011; Seo et al., 2011; Zhang et al., 2011). However, the mechanism by which BBSome deficiency causes changes in ciliary protein composition remains unclear.

Data from *Caenorhabditis elegans*, *C. reinhardtii*, and mammalian cells have revealed that only intact BBSomes enter cilia (Blacque et al., 2006; Lechtreck et al., 2009; Seo et al., 2010, 2011). Within cilia, BBSomes are moved by intraflagellar

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Abbreviations used in this paper: AMPK, AMP-regulated protein kinase; BBS, Bardet-Biedl syndrome; CAH6, carbonic anhydrase 6; DIC, differential interference contrast; GPCR, G protein-coupled receptor; IFT, intraflagellar transport; MCHR1, melanin-concentrating hormone receptor 1; PA, phosphatidic acid; PE, phosphatidylethanolamine; PLD, phospholipase D; SSTR3, somatostatin receptor 3.

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transport (IFT), a process by which large membrane-associated protein particles are transported bidirectionally along the axonemal microtubules (Kozminski et al., 1993; Rosenbaum and Witman, 2002; Blacque et al., 2004; Lechtreck et al., 2009). These particles consist of over 20 different proteins organized into two sub-complexes, termed IFT complexes A and B (Cole and Snell, 2009). Kinesin motors move these particles toward the ciliary tip (anterograde IFT); cytoplasmic dynein 1b (cytoplasmic dynein 2 in mammals) returns the particles to the cell body (retrograde IFT). Ciliary assembly is largely unaffected in *C. reinhardtii* *bbs* mutants and in *Bbs* knockout mice, and organization of IFT complexes A and B appears unaffected in the *C. reinhardtii* mutants (Mykityn et al., 2004; Lechtreck et al., 2009). BBS proteins are significantly less abundant than IFT proteins in cilia of *C. reinhardtii*, and only a subset of IFT particles carries BBosomes (Lechtreck et al., 2009). This suggests that the BBosome is an IFT cargo or cargo adapter dispensable for ciliary assembly and basic IFT in most cell types.

In mammalian cells, BBosome deficiencies affect the presence of ciliary transmembrane proteins, specifically of G protein-coupled receptors (GPCRs), in cilia. Melanin-concentrating hormone receptor 1 (MCHR1) and somatostatin receptor type 3 (SSTR3) fail to localize to neuronal cilia in *Bbs2*^{-/-} and *Bbs4*^{-/-} mice (Berbari et al., 2008b). The BBosome interacts with the ciliary targeting motif in the IP3 loop of SSTR3, and IP3^{SSTR3}-GFP fusion proteins translocate into cilia in a BBosome-dependent manner (Berbari et al., 2008a; Jin et al., 2010; Domire et al., 2011). The BBosome could facilitate the transport of proteins from the plasma membrane through the barrier of the ciliary transition zone into the ciliary membrane proper (Nachury et al., 2010). However, recent data show that the localization of some ciliary GPCRs is unaffected by a BBosome deficiency, whereas still others, e.g., dopamine receptor 1 in *Bbs4*^{-/-} mice and MCHR1 and the hedgehog effector Smoothened in *Bbs3*^{-/-} mice, accumulate in BBosome-deficient cilia (Domire et al., 2011; Seo et al., 2011; Zhang et al., 2011). This suggests that both import and export of ciliary GPCRs are affected in *Bbs* mutants.

We previously identified several putative signaling proteins (phospholipase D [PLD], an AMP-regulated protein kinase [AMPK], and a single domain globin [THB1]) that accumulate excessively in the ciliary membranes of *C. reinhardtii* *bbs1*, *bbs4*, and *bbs7* mutants (Lechtreck et al., 2009). Although lacking receptor functions, these membrane-associated proteins could modulate ciliary signaling, e.g., by protein phosphorylation and the synthesis of signaling lipids, providing a potential explanation for the disruption of phototactic behavior that is a hallmark of *bbs* mutants in *C. reinhardtii*.

To investigate the mechanism by which these signaling proteins are accumulated in the *bbs* cilia, we performed a detailed analysis of one of the proteins, PLD, in wild type versus the *bbs4-1* mutant that is null for BBS4 (Lechtreck et al., 2009). We chose to focus on PLD, both because it has a mammalian orthologue, PLD6, and because of its likely involvement in phospholipid signaling (Munnik et al., 2000). We have found that BBosome deficiency causes a massive redistribution of PLD from the cell body to the ciliary membrane, that the biochemical defects in *bbs* cilia increase with time but can be rapidly

corrected when wild-type BBosomes are introduced into the cytoplasm and cilia, that retrograde IFT acts upstream of the BBosome in the PLD export pathway, that PLD can enter the cilium independently of the BBosome and IFT, and, finally, that BBosome disruption causes secondary defects such as changes in the lipid composition of the ciliary membrane. We conclude that the *C. reinhardtii* BBosome functions downstream of IFT in the export phase of a process that cycles membrane signaling proteins through the cilium. The results further suggest that the absence of a protein from BBosome-deficient cilia is not necessarily due directly to failure of the BBosome to import the protein into the cilium.

Results

PLD redistributes from the cell body into cilia in *bbs* mutants

We previously reported, based on silver-stained gels, that PLD, AMPK, and THB1 are enriched in *C. reinhardtii* *bbs1*, *bbs4*, and *bbs7* cilia (Lechtreck et al., 2009). We have now generated an antibody that is specific for *C. reinhardtii* PLD (Fig. 1 A) and, using that antibody, are able to confirm that PLD is enriched in cilia of *bbs1*, *bbs4*, and *bbs7* cells (Fig. 1 B). In addition, we have extended the findings to a newly identified mutant for BBS8 (Fig. 1 B). This mutant has an insertion in the first exon of *BBS8* that results in a STOP codon predicted to terminate the normally 530-aa protein at aa 38 (Fig. S1 A). Therefore, this mutant is likely to be null for BBS8. Its isolated cilia lack BBS4, indicating that the BBosome without BBS8 is either unstable as in *bbs1-1* or is unable to enter the cilium as in *bbs7-1* (Lechtreck et al., 2009). To distinguish between these two possibilities, we assessed the level of BBS4 in the cytoplasm of the *bbs8-1* mutant; BBS4 was strongly reduced, indicating that the BBosome is largely unstable in the absence of BBS8 (Fig. S2 B). All four *bbs* mutants have normal length cilia (Fig. S2 A). Anterograde and retrograde IFT velocities as determined by differential interference contrast (DIC) microscopy are similar to that of wild type (Fig. S2 C); anterograde IFT frequency is also similar to wild type but retrograde IFT frequency appears to be slightly reduced (Fig. S2 D), although this was more difficult to measure because the retrograde particles are near the limit of detection for DIC. The levels of IFT particle proteins in the *bbs* cilia are similar to wild type (Fig. 1 C; and see Figs. 3 B and 6 A) or slightly elevated (Fig. S2 B and see Fig. 5 C); this slight increase, if real, may be an attempt by the mutant cells to compensate for loss of the BBosome. These results are consistent with our previous observation using immunofluorescence microscopy that the ciliary distribution of IFT complex A protein IFT139 and complex B protein IFT46 is unaffected in *bbs4-1* (Lechtreck et al., 2009).

Importantly, PLD can also be detected in wild-type cilia, particularly in the ciliary membrane, by means of Western blots probed with the anti-PLD antibody, indicating that PLD is a bona fide ciliary protein (Fig. 1, A and C). In wild type and *bbs4-1*, ciliary PLD is largely associated with the ciliary membrane, defined as the detergent phase of Triton X-114 phase partitioning (Fig. 1 C). To estimate just how great the

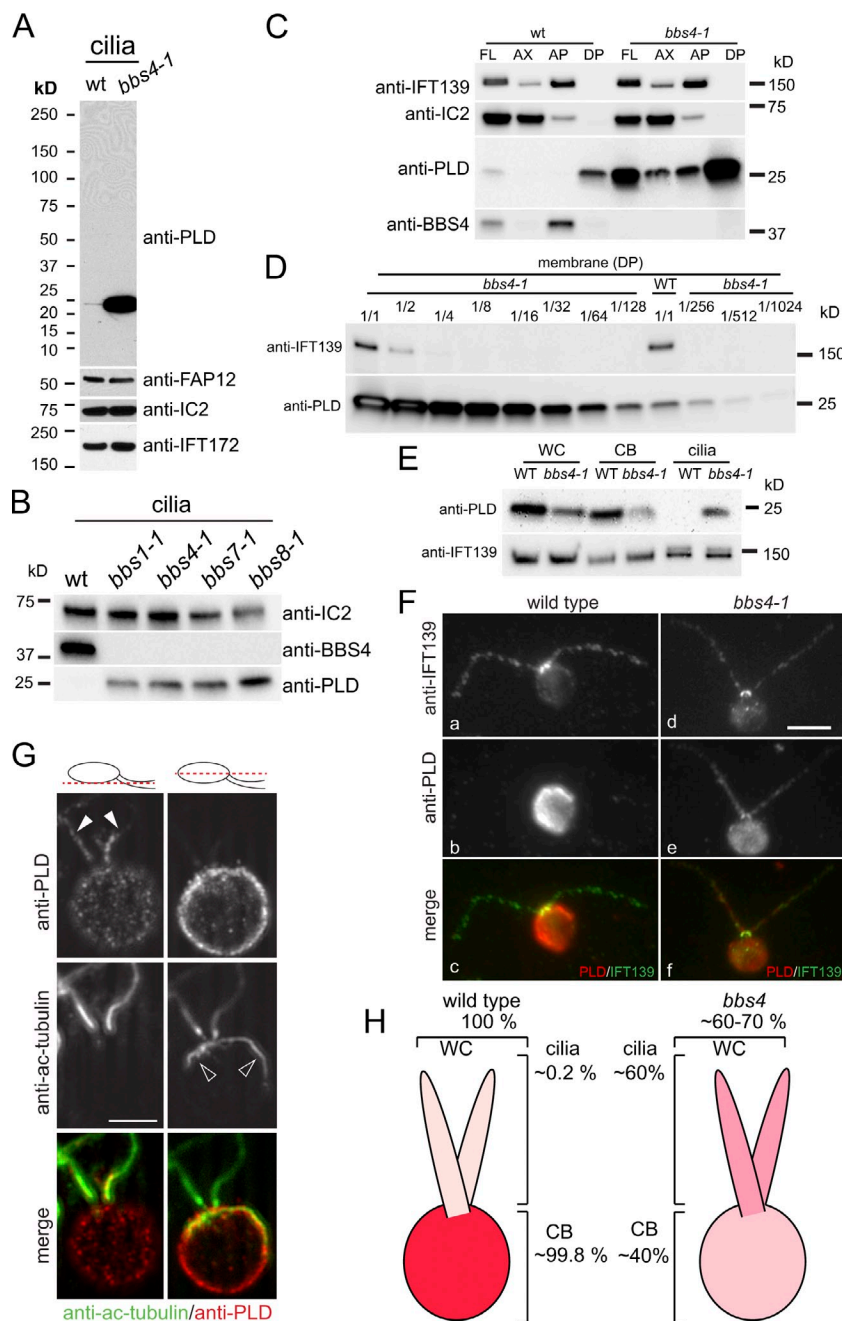


Figure 1. BBSome deficiency causes PLD redistribution from the cell body into cilia. (A) Western blot of isolated wild-type and *bbs4-1* cilia probed with antibodies to PLD and, as loading controls, antibodies to the membrane-associated protein FAP12, the axonemal dynein subunit IC2, and the IFT complex B protein IFT172. (B) Western blot of isolated cilia from wild type (wt) and *bbs1-1*, *bbs4-1*, *bbs7-1*, and *bbs8-1* probed with antibodies to BBS4 and PLD. IC2 was used as loading control. (C) Western blots of wild-type and *bbs4-1* cilia (FL) and ciliary fractions (AX, axonemes; AP and DP, aqueous and detergent phase from Triton X-114 phase partitioning, respectively) were probed with antibodies to PLD and BBS4, and with antibodies to the axonemal protein IC2 and the matrix protein IFT139 as loading controls. (D) Western blot of isolated ciliary membrane from wild type and *bbs4-1* probed with anti-PLD. The *bbs4-1* sample was diluted as indicated. The signal strength obtained from the undiluted wild-type sample is between that of the 1:128 and 1:256 dilution of the *bbs4-1* sample. (E) Western blots of whole cells (WC), deciliated cell bodies (CB), and isolated cilia (cilia) from wild type and *bbs4-1* were probed with anti-PLD. Equivalent amounts of cells and cilia were loaded (i.e., one whole cell, one cell body, and two cilia). The IFT particle protein IFT139 was used as a loading control. (F) Wild-type and *bbs4-1* gametes were stained with anti-IFT139 and anti-PLD. PLD was readily detected in cilia from *bbs4-1* but not detected in wild-type cilia. Bar, 5 μ m. (G) Two optical sections of a *bbs4-1* vegetative cell stained with anti-PLD and anti-acetylated tubulin. Merged images are shown in color. The positions of the optical sections are indicated in the diagrams. Solid arrowheads, cilia; open arrowheads, microtubular roots. Bar, 5 μ m. (H) Schematic presentation of PLD distribution in wild-type and *bbs4-1* mutant cells.

increase is in the mutant cilia, we used Western blotting to quantitate the relative amounts of PLD in isolated membranes of wild-type and *bbs4* mutant cilia. We determined that PLD is ~ 150 – $200\times$ more abundant in the mutant versus wild-type ciliary membrane (Fig. 1 D). A similar enrichment was observed in cilia of mutant versus wild-type gametes (unpublished data).

To distinguish whether the observed enrichment of PLD in the cilia of *bbs* mutants is a result of a general increase in PLD levels in the mutant cells or to defects in ciliary trafficking of PLD, we determined the distribution of PLD in wild-type and *bbs* mutant cells and cilia. Western blotting revealed that the total amount of PLD in *bbs4-1* cells actually was reduced by $\sim 40\%$ in comparison to wild-type cells (Fig. 1, E and H). Moreover, in

wild type, $\sim 0.2\%$ of the total PLD is in the cilia, whereas in the mutant, $\sim 60\%$ of the total PLD is in the cilia.

Immunofluorescence microscopy confirmed our biochemical data: PLD was readily detected in a spotted distribution along the length of *bbs4-1* mutant cilia but was barely detectable in wild-type cilia (Fig. 1 F and see Fig. 4 B). Within mutant cilia the signal strength was more pronounced in the proximal region of the cilium. Labeling by the anti-PLD antibody was similar in wild-type and *bbs4-1* cell bodies, albeit signal strength was reduced in the mutant. Label was distributed in a spotted pattern at the cell periphery (Fig. 1 G for *bbs4-1*); this could represent vesicles underlying the plasma membrane. In summary, the *bbs4-1* mutation causes a dramatic redistribution of PLD from the cell body into cilia (Fig. 1 H).

Table 1. Quantitative analysis of phospholipids and DAG in wild-type and *bbs4-1* cilia

Phospholipid	Total phospholipid in wild-type cilia	Total phospholipid in <i>bbs4-1</i> cilia	Change in <i>bbs4-1</i> cilia
	%	%	%
DAG 18:1/18:1	1.35	1.95	+44
DAG 18:3/18:0	1.55	9.29	+500
PA 18:3/18:0	11.36	16.84	+50
PE 18:0/18:3	49.64	35.12	-29
PE 18:1/18:3	1.77	1	-43
PG 16:0/16:0	20	22.6	+13
PG 16:1/16:1	3	3.74	+25
PG 18:0/16:0	0.46	0.46	+1
PG 18:1/16:0	8.68	7.3	-16
PI 18:1/16:0	2.89	1.7	-41

The distribution of phospholipids in cilia of wild type and *bbs4-1* as a percentage of total phospholipid. PG, phosphatidylglycerol; PI, phosphatidylinositol.

The lipid composition of *bbs4* cilia is altered

To assess whether the accumulation of PLD in *bbs* cilia affects the lipid composition of the ciliary membrane, we performed a metabolomic and lipidomic analysis of purified wild-type and mutant cilia (Table 1 and Fig. 2). Phosphatidylethanolamine (PE; Fig. 2, A and B), the substrate of PLD in *C. reinhardtii* (Munnik et al., 2000), was reduced by ~30–50% in *bbs4-1* cilia. Phosphatidic acid (PA) and ethanolamine, the products of PLD activity, were increased by 50% and 4,000%, respectively (Table 1 and Fig. 2 C). DAG and stearic acid, putative downstream products of PA enzymatic degradation, were each increased by ~500%; a two- to threefold increase was also observed for glycerol and glycerol phosphate (Fig. 2 D and not depicted). The changes in PE, PA, and DAG occur predominately in species carrying 18:3/18:0 fatty acids, indicating that they are linked, i.e., that DAG 18:3/18:0 is formed by dephosphorylation of PA 18:3/18:0 (Table 1). The amounts of other membrane lipids, including that of the major ciliary membrane lipids ergosterol and stigmaterol, which account for ~32% and ~20% of the total ciliary lipid, respectively, were similar in both strains (unpublished data). In conclusion, the lipid composition of the ciliary membrane of the *bbs4-1* mutant is altered and the observed changes are as predicted for increased PLD activity. The data also reveal that only a limited number of phospholipid species are present in the ciliary membrane of *C. reinhardtii* (Table 1).

The defects in protein content of *bbs4* cilia increase with time

To distinguish whether the abnormal amounts of PLD in *bbs* cilia are deposited there during initial ciliary assembly or accumulate there with time after ciliary formation, we took advantage of the ability of *C. reinhardtii* to fully assemble cilia de novo within ~60 min after amputation of the cilia by pH shock (Fig. 3 and Fig. S3 E). We simultaneously deciliated large populations of wild-type and *bbs4-1* cells and allowed them to synchronously regenerate their cilia; we then isolated the recently formed cilia at time points ranging from 80 to 380 min after deciliation and compared them by SDS-PAGE and Western blotting (Fig. 3 A). At the loadings used, PLD was not detected in wild-type whole cilia (Fig. 3 B, top), but small amounts of

PLD were consistently detected in isolated wild-type ciliary membranes, where the PLD was maintained at a relatively constant low level during the time period analyzed (Fig. 3 B, bottom). In contrast, PLD was detectable in the newly assembled mutant whole cilia at least as early as 80 min after deciliation; it then accumulated steadily over the course of several hours as the cilia aged (Fig. 3, B [top] and C). The data indicate that PLD accumulation in mutant cilia is a dynamic process that continues in full-length steady-state cilia. The amounts and kinetics of accumulation of several ciliary transmembrane proteins, specifically PKD2, FMG1, and the mastigoneme protein, were similar in wild-type and mutant cilia, suggesting that the transport of these proteins is unaffected by BBS4 deficiency.

In the course of the cell cycle, cilia are normally formed after division, at least 6 h before our synchronously grown cells were used for biochemical analyses. We here term such 6-hour-old cilia “standard cilia.” Comparison of regenerated and standard cilia from wild type and *bbs4-1* using silver-stained gels revealed a protein of ~24 kD that was present in wild-type cilia and in newly regenerated *bbs4* cilia but was significantly reduced in standard *bbs4* cilia (Fig. 3 D). The protein was tentatively identified as carbonic anhydrase 6 (CAH6) based on recovery of a single peptide by mass spectrometry (Fig. S1 B); the identification was confirmed using an antibody to *C. reinhardtii* CAH6 (Fig. 3 B). Quantitative analysis of age-sorted cilia showed that at 80 min after deciliation the mutant cilia contained ~50% of the CAH6 present in comparable wild-type samples (Fig. 3, B and E). Over the next five hours CAH6 was progressively lost from mutant cilia, whereas little reduction of CAH6 was observed in wild-type cilia (Fig. 3, B, D, and E). Because a substantial amount of CAH6 is present in newly formed mutant cilia, it is unlikely that BBSome defects inhibit CAH6 ciliary import. The data rather suggest that *bbs* cilia increasingly lose the ability to retain CAH6. Thus, CAH6 loss from *bbs* cilia appears to be caused indirectly by the BBSome defect.

Excess PLD is rapidly removed from *bbs* cilia when BBSomes are reintroduced

The progressive redistribution of PLD from the cell body into cilia of *bbs* mutants could be caused by a failure of PLD ciliary export. Therefore, to test directly whether BBSomes

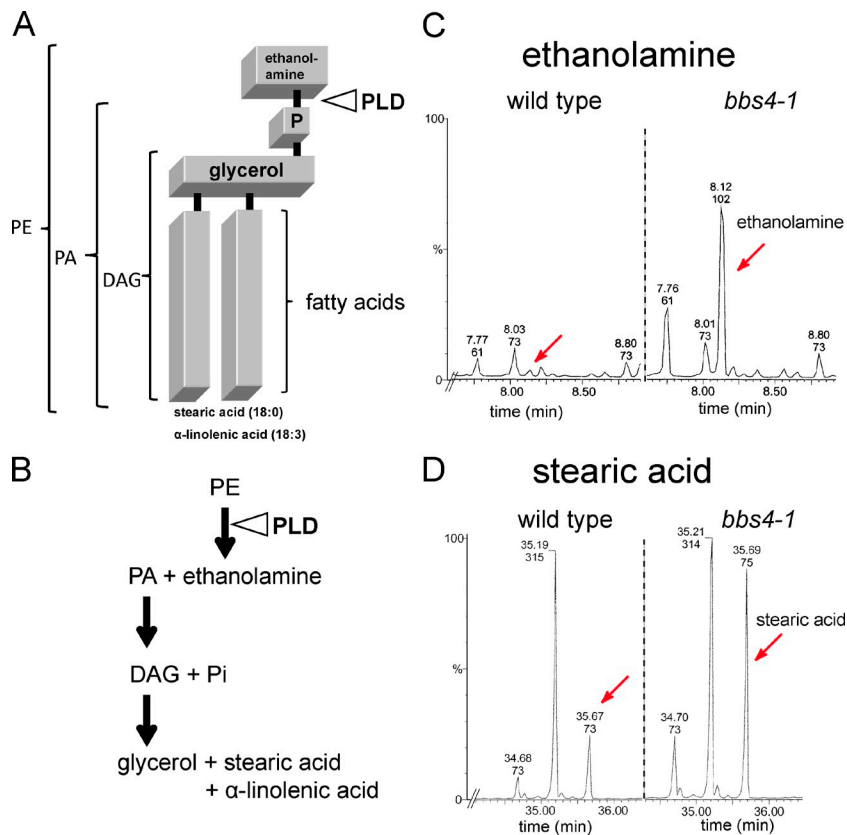


Figure 2. The lipid composition of *bbs4-1* cilia is altered. (A) Schematic presentation of the PLD substrate PE 18:0/18:3 and the site of PLD cleavage. (B) Schematic presentation of the enzymatic breakdown of PE into PA, ethanolamine, DAG, and downstream products. (C and D) Gas chromatography-mass spectrometry profiles showing the elution times for small metabolites from wild-type and *bbs4-1* cilia. Ethanolamine (peaks marked by red arrows in C) is enriched $\sim 40\times$ and free stearic acid (peaks marked by red arrows in D) is enriched $\sim 6\times$ in *bbs4-1* cilia. Accuracy of quantitation is $\pm 10\%$.

are required to remove PLD from cilia, *bbs4* and wild-type gametes were mixed and the distribution of PLD in cilia of the resulting zygotes was analyzed at various time points after mating. The zygotes possess four cilia, two derived from the mutant gametes and two from the wild-type gametes (Fig. 4 A). After cell fusion, BBSomes contributed by the wild-type cytoplasm become immediately available to *bbs4* mutant cilia. With immunofluorescence microscopy, PLD is readily detectable in cilia of mutant but not wild-type gametes (Fig. 4 B, a–c). Ten minutes after mixing of the gametes, some zygotes were observed that possessed two cilia with PLD derived from the *bbs4* parent and two cilia without detectable PLD derived from the wild-type parent (Fig. 4, B [d–f] and C). However, even at this early time point, PLD was reduced or not detectable in cilia of the majority of zygotes (Fig. 4, B [g–i] and C). The share of zygotes without detectable ciliary PLD increased progressively from 18% at 10 min to 96% at 50 min after mixing of the gametes (Fig. 4 C). The time span required for gametes to complete fusion after mixing is variable—some zygotes may form as early as 15 s after mixing, with typically 50% of the gametes having completed fusion after 5 min in good mating reactions (Hunnicut and Snell, 1991). The presence of a few zygotes with cilia still containing PLD at later time points can be explained by delayed fusion of some gametes. In conclusion, PLD is removed in <10 min from the formerly mutant cilia once BBSomes become available. Therefore, an efficient pathway for PLD export from cilia is present in *C. reinhardtii* and this pathway requires the BBSome.

PLD export from cilia requires retrograde IFT

We previously showed that the BBSome cycles in association with IFT through cilia (Lechtreck et al., 2009). To determine if this association is important for BBSome function and normal export of PLD from the cilium, we analyzed the mutant strain *dhc1b-2* (previously referred to as *dhc1b^{ts}*). This strain contains a hypomorphic allele of the gene encoding the heavy chain of dynein 1b, the motor for retrograde IFT (Lechtreck et al., 2009; Witman, 2012). At the permissive temperature (21°C) used for all experiments involving *dhc1b-2* here, $\sim 60\%$ of the mutant cells assembled two cilia that were slightly shorter than those of wild type (Fig. S3, A and B). The velocity and frequency of retrograde IFT and the amounts of the retrograde dynein subunits DHC1b and D1bLIC were reduced in the *dhc1b-2* cilia (Fig. 5 A and Fig. S2, C and D). The amount of BBS4 was elevated in *dhc1b-2* cilia, as expected for a protein cycled through the cilium by IFT. However, despite the presence of BBS4/BBSomes, PLD was highly enriched in the mutant cilia (Fig. 5 A), indicating that the BBSome-dependent export of PLD from cilia requires the retrograde motor for IFT.

As an independent test of the involvement of IFT in export of PLD, we analyzed the mutant *ift74-1*, which expresses a truncated version of the IFT complex B protein IFT74 (Fig. 5 B; unpublished data). The truncated IFT74 is assembled into complex B and transported into the cilium as part of the IFT particle; the ciliary levels of the other complex B proteins are near normal, but the amount of IFT complex A is greatly reduced. Compared with wild-type cilia, BBS4 was reduced by $\sim 65\%$ and

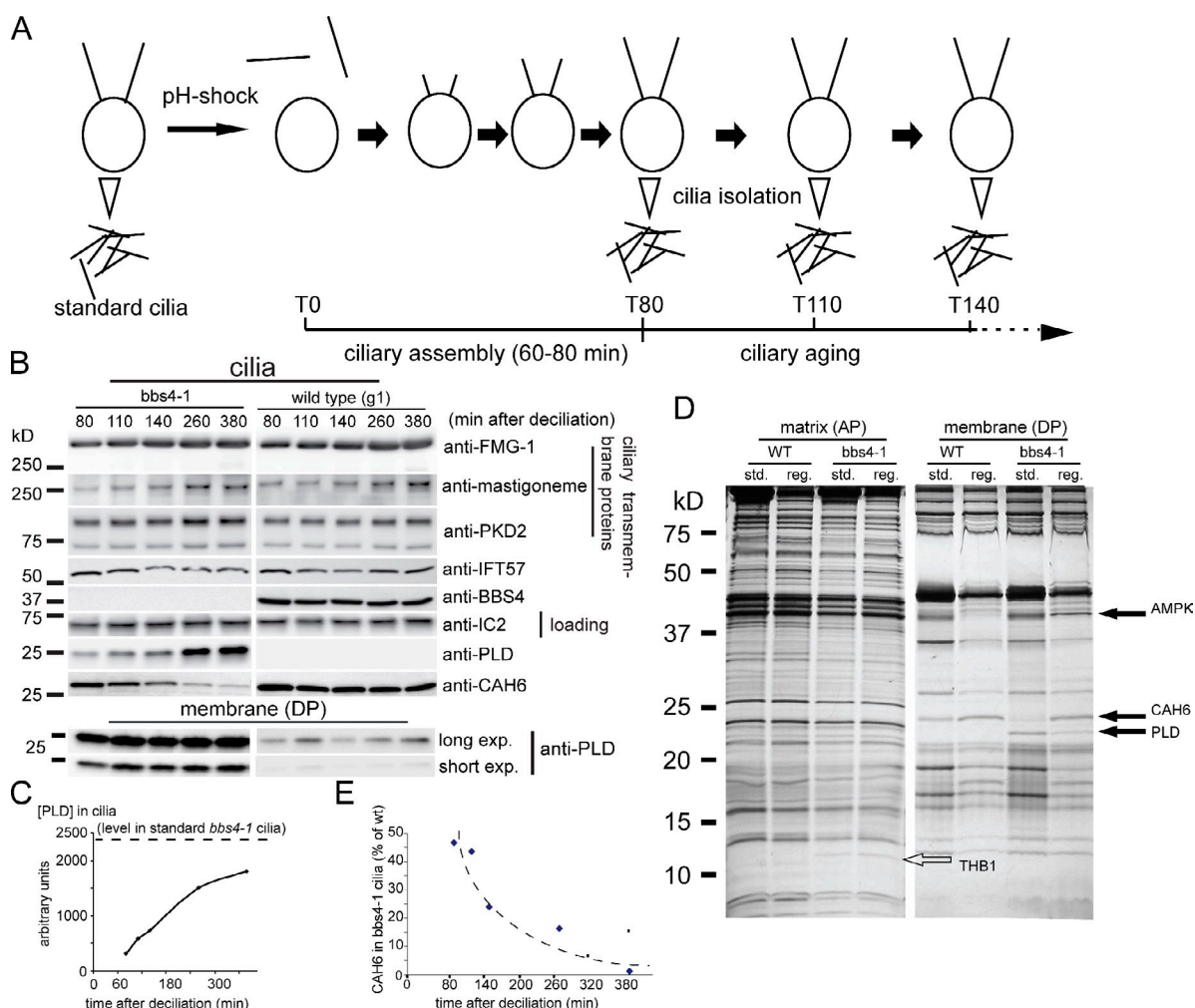


Figure 3. The biochemical defects in *bbs4-1* cilia increase with time. (A) Schematic presentation of the experimental design for obtaining age-sorted cilia from *C. reinhardtii*. (B) Western blots comparing regenerated wild-type and *bbs4-1* cilia isolated at various time points after deciliation. (top) Western blots were probed with anti-FMG-1, anti-mastigoneme protein, anti-PKD2, anti-IFT57, anti-BBS4, anti-IC2 as a loading control, anti-PLD, and anti-CAH6. Ciliary length measurements for a similar experiment are shown in [Fig. S3 E](#). (bottom) Long and short exposures of a Western blot of isolated ciliary membrane fractions from the same experiment probed with anti-PLD. (C) Graph showing the accumulation of PLD with time in *bbs4-1* cilia; the dashed line corresponds to PLD levels in standard *bbs4-1* cilia, i.e., cilia harvested without prior deciliation at least 6 h after the beginning of the light phase. (D) Silver-stained gel comparing matrix and membrane isolated from standard and 2-h-old regenerated wild-type and *bbs4-1* cilia. Solid arrows, proteins altered in *bbs4-1* ciliary membrane; open arrow, THB1. Note the differences in composition in the membrane of young and old cilia; no major differences were observed between young and old axonemal (not depicted) and matrix fractions, indicating that age-induced changes in ciliary protein composition are largely restricted to the ciliary membrane. (E) Graph showing the loss of CAH6 in *bbs4-1* cilia as a percentage of that in wild-type cilia over time. The data shown in C and E are from a single representative experiment out of two repeats.

PLD was strongly accumulated in *ift74-1* cilia (Fig. 5 B). The data suggest that entry of BBSomes into cilia requires IFT74 and/or IFT complex A and that the residual IFT and BBSomes in *ift74-1* cilia are insufficient to remove PLD from cilia. The result is consistent with a recent observation in *C. elegans* that a mutation in complex A protein IFT144 abrogates entry of BBSomes into cilia (Wei et al., 2012).

IFT complex A has an important role in retrograde IFT (Pedersen and Rosenbaum, 2008; Pedersen and Christensen, 2012). To more directly investigate if complex A has a role in BBSome transport and PLD export, we analyzed strains *fla15-1* and *fla17-1*, which carry mutations in the complex A proteins IFT144 and IFT139, respectively (Piperno et al., 1998; Iomini et al., 2009). The *fla15-1* strain has a point mutation that converts a highly conserved cysteine in IFT144 to an arginine.

The *fla17-1* strain has an in-frame deletion of three exons of the *IFT139* gene and produces a smaller-than-normal version of IFT139 (Fig. 5 C; Iomini et al., 2009). These *fla* mutants fail to assemble cilia at elevated temperatures. At the permissive temperature used here IFT is sufficient for ciliary assembly but the velocity and frequency of retrograde IFT is reduced and the cilia develop small bulges containing complex B components, indicative of a defect in returning cargo to the cell body (Iomini et al., 2009). Compared with wild type, BBS4 was highly enriched in cilia of *fla17-1* (Fig. 5 C), suggesting that the BBSome also is not returned normally to the mutant cell body. Despite the abundance of BBSomes, PLD also was highly enriched in *fla17-1* cilia (Fig. 5 C), indicating that PLD export from cilia is dependent on IFT complex A. In contrast, the amounts of BBS4 and PLD were near normal in

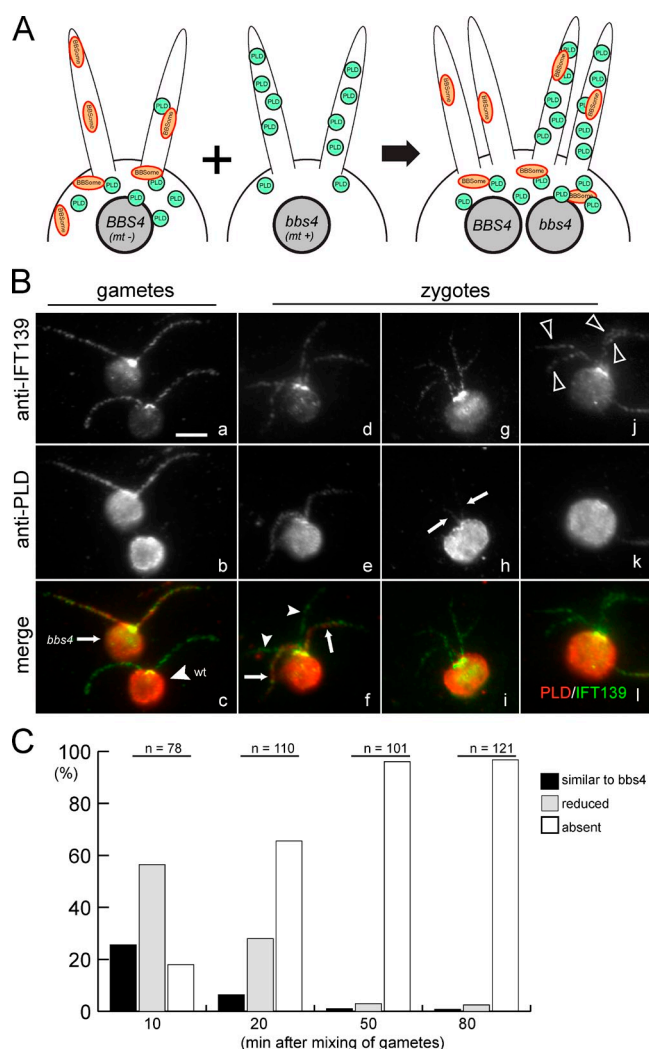


Figure 4. PLD is removed from cilia of *bbs4-1* cells when BBSomes are introduced by fusion with wild-type cells. (A) Schematic presentation of the PLD ciliary export assay. In wild-type gametes BBSomes are present in both cytoplasm and cilia, and nearly all of the cell's PLD is in the cell body; *bbs4-1* gametes lack BBSomes and have undergone a massive redistribution of PLD from the cell body to the cilia. Wild-type (*BBS4*, *mt*⁻) and *bbs4-1* (*bbs4*, *mt*⁺) gametes were mixed and allowed to fuse. The resulting zygotes have four cilia (two each from the wild-type and mutant gametes); BBSomes are present in the shared cytoplasm. (B) Gametes (a–c) and quadriliated zygotes (d–l) were stained with antibodies to PLD and to IFT139 to visualize the cilia. Arrow in c, *bbs4* gamete; arrowhead in c, wild-type gamete; arrows in f and h, cilia containing PLD derived from mutant gametes; arrowheads in f, cilia lacking PLD derived from a wild-type gamete; arrowheads in j, four cilia of a zygote without detectable PLD. Bar, 5 μ m. (C) Quadriliated zygotes were scored by eye for the amount of PLD in their cilia. Black bars, quadriliated cells with two wild-type cilia and two cilia with levels of PLD characteristic of the *bbs4-1* gamete; gray bars, quadriliated cells with two wild-type cilia and two cilia with detectable but reduced amounts of PLD; white bars, quadriliated cells without detectable PLD in their cilia.

cilia of *fla15-1* (Fig. 5 C). The data suggest that IFT139 is involved in retrograde transport of BBSomes and removal of PLD from cilia.

PLD can enter cilia independently of IFT

Our data suggest that retrograde IFT particles transport PLD in a BBSome-dependent manner. Because PLD enters cilia through a BBSome-independent process, we investigated if IFT is required

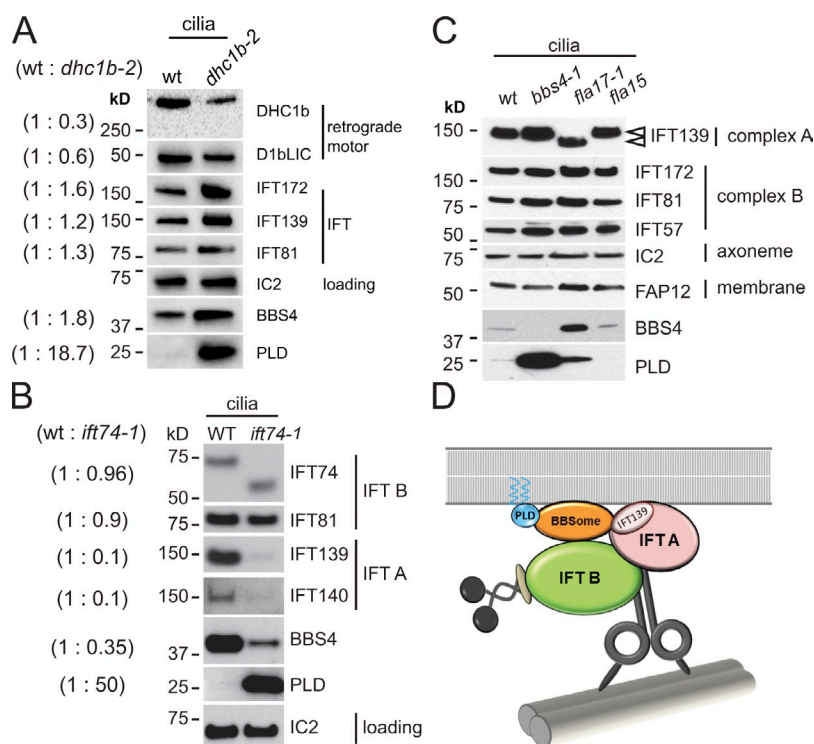
for entry of PLD into cilia. We took advantage of the *fla10-1* mutant, which carries a temperature-sensitive mutation in one of the heavy chains of the anterograde IFT motor kinesin-2, so that IFT can be rapidly turned off by a temperature shift (Kozminski et al., 1995). At the permissive temperature (22°C), *fla10-1* cilia contained slightly elevated amounts of PLD, which can be explained by the somewhat reduced amounts of IFT complex A and BBSomes (as indicated by reduced amounts of IFT139 and BBS4, respectively) in the mutant cilia (Fig. 6 A). Cilia isolated from *fla10-1* that had been incubated for 2.5 h at the nonpermissive temperature (32°C) were largely depleted of IFT81 (complex B) and IFT139 (complex A). This loss of IFT proteins at the restrictive temperature was accompanied by an increase in ciliary PLD; in contrast, the amounts of PLD in wild-type and *bbs4-1* cilia were essentially unaffected by the temperature shift. To determine the temporal sequence of IFT depletion and PLD accumulation in *fla10-1* cilia, we analyzed cilia isolated at various time points (1–6 h) after shifting *fla10-1* cells to 32°C (Fig. 6 B). During the incubation time the number of mutant cells carrying one or two cilia decreased from >90% to ~70% and the mean length of the cilia that they did have decreased from ~11 to ~6 μ m; the number and length of cilia of wild type was unaffected at 32°C (Fig. S3, C and D). Starting at ~2 h at 32°C, by which time the level of IFT proteins was considerably reduced, PLD began to accumulate in *fla10-1* cilia. Importantly, PLD continued to accumulate in *fla10-1* cilia after 4 h at 32°C when IFT proteins were nearly undetectable; after ~6 h at 32°C it reached a level approaching that of *bbs* mutants (>30 $\times$ above the level before temperature shift; Fig. 6 C). The data indicate that PLD entry into cilia does not require IFT. Note that despite the depletion of IFT proteins, BBS4 remained present in *fla10-1* cilia. The accumulation of PLD in the presence of BBS4 provides additional evidence that BBSomes alone are insufficient to control PLD levels in cilia in the absence of IFT.

Discussion

BBSome deficiency has been shown to alter the membrane protein composition of cilia in *C. reinhardtii* and mammals (Berbari et al., 2008b; Lechtreck et al., 2009; Domire et al., 2011; Seo et al., 2011; Zhang et al., 2011). We showed previously that BBSomes travel through cilia in association with a subset of IFT particles (Lechtreck et al., 2009). These observations raise the possibility that BBSomes contribute to IFT-mediated transport of ciliary membrane proteins. Here we analyzed the role of the BBSome and IFT in the ciliary transport of PLD, a membrane-associated protein that accumulates in cilia of *C. reinhardtii* *bbs* mutants. Our data reveal that PLD continuously enters the cilium in a BBSome- and IFT-independent manner but that export of PLD from the cilium is dependent on the BBSome, probably functioning as a cargo adaptor for IFT.

PLD cycles through the cilium and redistributes from the cell body to the cilium in the absence of the BBSome or IFT
PLD is abundant in wild-type cell bodies and ~0.2% of the PLD is present in wild-type cilia. However, in *bbs4-1* mutants

Figure 5. Ciliary PLD is elevated in retrograde IFT mutants. (A) Western blot of isolated wild-type and *dhc1b-2* cilia probed with antibodies to the proteins indicated. The amounts of the dynein 1b heavy chain, DHC1b, and light intermediate chain, D1bLIC, are reduced in the mutant cilia because of the instability of the dynein motor complex. BBS4 and in particular PLD are enriched in mutant cilia. IC2 was used as a loading control. For quantification of relative protein levels, values for band intensities were first normalized to the IC2 band intensity and the ratios (left, parentheses) were then calculated. (B) Western blot of cilia isolated from wild type and *ift74-1* probed with antibodies to the proteins indicated. In *ift74-1* the IFT complex B protein IFT74 is truncated. Note reduced amounts of IFT complex A proteins and BBS4 and increased levels of PLD in the mutant cilia. Relative protein levels (in parentheses) were determined as in A. (C) Western blot probing isolated cilia from wild type, *bbs4-1*, *fla17-1* (expressing a truncated IFT139; arrowheads), and *fla15-1*. Note increase of BBS4 and PLD in *fla17-1*. (D) Summary diagram showing the IFT motors, IFT complexes A and B, the BBSome, and PLD. The BBSome functions as an IFT cargo adapter linking PLD and other cargoes to retrograde IFT particles. PLD is predicted to be myristoylated and palmitoylated (Fig. S4 C), offering a possible explanation for its membrane association. Dynein is shown linking the IFT particle to a doublet microtubule, which provides a track for IFT.



there is a massive redistribution of PLD from the cell body into the cilia, such that about half of the PLD is now contained in the cilia. Taking into account the relative volumes of the cell body ($>500 \mu\text{m}^3$) and the cilium ($<0.5 \mu\text{m}^3$; and ignoring possible differences in the fraction of each that is accessible to PLD), PLD in *bbs4-1* cells is enriched $\sim 1,000\times$ in the two cilia relative to the cell body. Because this concentration occurs in *bbs* mutants, PLD must be able to move into the cilia in the absence of BBSomes. A similar great enrichment was observed in cilia of the conditional mutant *fla10-1* after IFT proteins (but not BBS4) were depleted by incubation at the nonpermissive temperature. This strongly suggests that entry of PLD into cilia can occur even in the absence of IFT. Recently, Kee et al. (2012) reported that small molecules and proteins ≤ 40 kD can freely pass from the cytoplasm into cilia. Thus, PLD, which is only 25 kD, may enter the cilia slowly by diffusion and then become trapped there in the absence of an active export mechanism.

Because PLD accumulates progressively over time in BBSome- and IFT-deficient cilia, it is likely that the protein is continuously cycled through the cilium in wild-type cells, with IFT-dependent export mediated by the BBSome. The high efficiency of this PLD export pathway, as demonstrated by the rapid clearing of accumulated PLD when BBSomes are reintroduced into *bbs* mutant cilia, suggests that PLD entering wild-type cilia will be rapidly picked up and removed from the organelle. A system in which the movement of a protein into the cilium is BBSome independent but its export is BBSome dependent would allow fine tuning of the amount of the protein in the cilium by regulating the number of BBSomes that enter the cilium or by regulating the affinity of the BBSome for the protein.

A similar role for the BBSome in mammalian primary cilia is suggested by recent observations on dopamine receptor 1, a GPCR that is rapidly transported to and from cilia in response to environmental cues (Domire et al., 2011). Import of dopamine receptor 1 into cilia is BBSome independent whereas translocation of the protein from cilia requires the BBSome (Domire et al., 2011; Zhang et al., 2011). In addition, very recent results indicate that accumulation of the hedgehog signaling proteins Smoothened and Patched 1 occur in cilia of cells from *Bbs^{-/-}* mice (Zhang et al., 2012). Like PLD in *C. reinhardtii*, Smoothened is believed to continuously shuttle through the cilium (Ocbina and Anderson, 2008). Thus, BBSome-mediated export may be a feature of many membrane-associated proteins that undergo such cycling.

BBSome function depends on retrograde IFT

To investigate the role of IFT in the BBSome-dependent export of PLD from the cilium, we analyzed the distribution of BBS4 and PLD in cilia isolated from the dynein 1b mutant *dhc1b-2*, which is characterized by reduced retrograde movement of IFT particles. Using silver-stained 1D and 2D gels, we previously determined that THB1 is enriched in *dhc1b-2* cilia, but we could not ascertain the levels of PLD in these cilia (Lechtreck et al., 2009). Now, using our anti-PLD antibody and Western blotting, we found that PLD also is significantly accumulated in *dhc1b-2* cilia despite the presence of BBSomes. Likewise, BBSomes are present albeit reduced during PLD accumulation in *fla10-1* cilia at restrictive temperature. Therefore, the presence of BBSomes in cilia is insufficient to remove PLD when IFT is defective. Because the primary effect of the *dhc1b-2* mutation is on retrograde IFT, these results indicate that retrograde IFT is critical

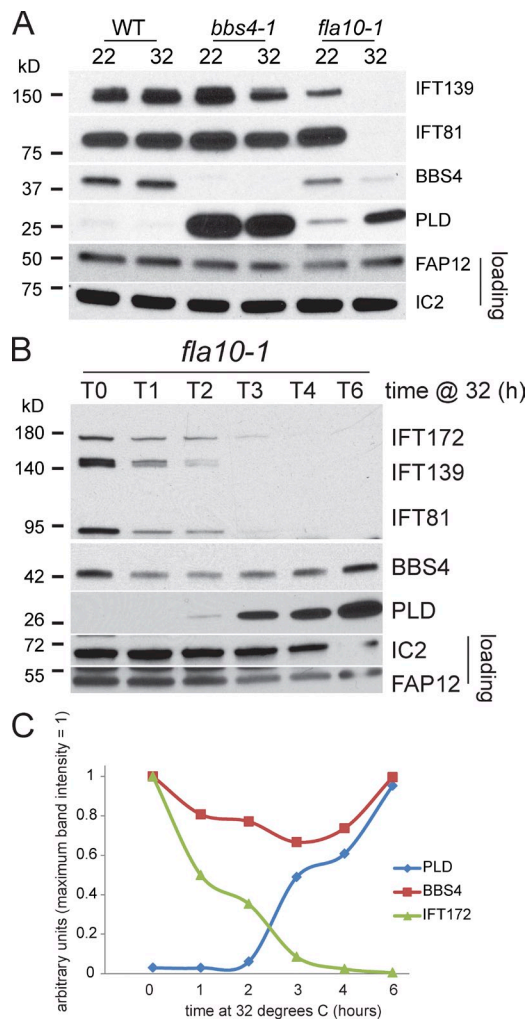


Figure 6. PLD accumulates in cilia after depletion of IFT proteins. (A) Western blot of cilia isolated from wild type, *bbs4-1*, and *fla10-1* at 22°C and after 2.5 h at 32°C. The blot was probed with antibodies to the proteins indicated. Depletion of IFT proteins and accumulation of PLD in response to the temperature shift was observed only in cilia from *fla10-1*. (B) Western blot comparing *fla10-1* cilia isolated at various time points from 0 to 6 h (T0–T6) after shifting the cells to 32°C. The blot was probed with antibodies to the proteins indicated. Note progressive depletion of IFT proteins and concomitant accumulation of PLD. (C) Relative amounts of PLD, BBS4, and IFT172 in the Western blot analysis shown in B. Maximum band intensity for each protein was set to 1. At later time points the mean length of *fla10-1* cilia was reduced (Fig. S3 D). Because similar amounts of protein were loaded for each sample, the slight increase observed for BBS4 during the second half of the incubation time is likely to reflect the higher number of cilia loaded.

for PLD export from the cilium. Inasmuch as loss of the BBSome has little if any effect on IFT (Figs. 1 C, 3 B, 6 A, and S2; Lechtreck et al., 2009) but defects in retrograde IFT affect export of both the BBSome and PLD, IFT must function upstream of the BBSome in the PLD export pathway; i.e., retrograde IFT moves the BBSome out of the cilium, and the BBSome in turn carries PLD from the cilium (Fig. 5 D).

Export of Smoothed from mammalian primary cilia similarly is dependent on the retrograde IFT motor, as indicated by the accumulation of Smoothed in cilia of mouse cells having a mutation in the dynein 2 heavy chain *Dync2h1* (Ocbina

and Anderson, 2008) or in which the same protein was knocked down by shRNA (Kim et al., 2009) or inhibited pharmacologically (Firestone et al., 2012). Thus, IFT-dependent BBSome-mediated export of signaling molecules from the cilium may be widespread. However, another hedgehog signaling protein, Gli2, which also is believed to cycle through the cilium (Kim et al., 2009), does not accumulate in *Bbs*^{−/−} mice (Zhang et al., 2012), suggesting the existence of distinct export (and import) mechanisms for different ciliary proteins.

IFT complex A is critical for BBSome export

The aforementioned observations suggest that coupling of PLD to retrograde IFT particles requires the BBSome. To explore how the BBSome interacts with the IFT machinery, we used mutants defective in IFT particle proteins. IFT complex A proteins and BBS4 are similarly decreased in cilia of *ift74-1*, which expresses a truncated complex B IFT74 protein. PLD is highly enriched in *ift74-1* cilia, indicating that the residual BBSomes are insufficient for its export. This suggests that IFT74 is critical for the entry of IFT complex A into cilia and that complex A could mediate binding of the BBSome to IFT complex B and the associated dynein motors during retrograde IFT. To test for a possible role of IFT complex A as a docking site for the BBSome, we analyzed *fla15* and *fla17-1*, which are defective in the IFT complex A proteins IFT144 and IFT139, respectively, and have reduced retrograde IFT (Piperno et al., 1998; Iomini et al., 2009). The distributions of BBS4 and PLD were nearly normal in *fla15*, whereas PLD and in particular BBS4 were enriched in cilia of *fla17-1*. Therefore, IFT139 could be involved in binding the BBSome to IFT complex A during retrograde IFT, rendering it critical for the export of the BBSome and PLD from cilia (Fig. 5 D). It may be relevant that the Tubby family protein TULP3 similarly interacts with complex A in mammals to promote ciliary trafficking of a subset of GPCRs, including MCHR1, and that retrograde transport of TULP3 specifically requires IFT139 (Mukhopadhyay et al., 2010).

In summary, PLD accumulates in cilia of mutants with defective BBSomes (*bbs4-1*, *bbs1-1*, *bbs7-1*, and *bbs8-1*; Lechtreck et al., 2009), with reduced amounts of BBSomes in cilia (*ift74-1*), and with reduced retrograde IFT of the BBSome (*dhc1b-2* and *fla17-1*). We propose that the BBSome functions as a cargo adapter linking PLD to retrograde IFT particles to ensure export of PLD from cilia. Further work will be necessary to understand how PLD is delivered to the cilium, how PLD interacts with the BBSome during the export phase of its cycle, and how the BBSome interacts with IFT complexes A and B.

BBSome function in IFT assembly and turnaround

In *C. elegans* mutants null for *bbs-7* and *bbs-8*, IFT complex A separates from complex B and the two complexes are transported anterogradely by kinesin-2 and OSM-3, respectively (Ou et al., 2005). These observations led to the hypothesis that the BBSome couples complex A to complex B and coordinates the activity of the two anterograde IFT motors. However, more recently it has been reported that a hypomorphic mutation at a

conserved site in the complex A protein DYF-2 (the *C. elegans* homologue of IFT144) causes complete dissociation between the BBSome and moving IFT particles, yet complexes A and B remain associated in anterograde IFT (Wei et al., 2012). These results indicate that in *C. elegans* the BBSome is not essential for stabilizing the binding of complex A to complex B, in good agreement with our previous observation that the colocalization of complexes A and B is not affected in cilia of *C. reinhardtii* *bbs4-1* mutants (Lehtreck et al., 2009). In the *C. elegans* IFT144 mutant, the BBSome is still formed but remains at the base of the cilium. Moreover, complex B but not complex A accumulates at the tip of the cilium, suggesting that complex B fails to be coupled to complex A for retrograde IFT. Based on these observations, Wei et al. (2012) proposed that in wild-type *C. elegans* the BBSome regulates assembly of complexes A and B into intact IFT particles at the base of the cilium, and is then carried as cargo by IFT to the tip of the cilium, where it has a similar role in reorganizing the IFT particle for retrograde transport. In the mutant, the BBSome still functions to assemble the IFT particle at the base of the cilium, but because it is no longer transported to the ciliary tip, IFT particle reorganization is defective and complex B accumulates there.

We previously reported that in *C. reinhardtii* the BBSome is not an integral part of the IFT machinery but rather appears to be a cargo or cargo adapter, a conclusion now supported by the observations of Wei et al. (2012). However, in *C. reinhardtii* loss of the BBSome does not seem to affect the assembly of complexes A and B (Lehtreck et al., 2009), does not lead to an accumulation of complex B at the tip of the cilium (Lehtreck et al., 2009), and does not obviously alter the balance of complex A to complex B within the cilium as would be expected if complex B were being left behind during retrograde transport (Figs. 5 C, 6 A, and S2 B). Therefore, in contrast to the situation in *C. elegans*, it is unlikely that the BBSome has a key role in regulating IFT particle assembly at either the base or the tip of the *C. reinhardtii* cilium.

The function of PLD in *C. reinhardtii*

The function of PLD in *C. reinhardtii* is likely to involve phospholipid signaling (Meijer et al., 2002). Consistent with this, we found that the accumulation of PLD in cilia of *C. reinhardtii* mutants is correlated with major changes in the phospholipid content of the ciliary membrane. It is possible that PLD cycles through the wild-type cilium as part of a system that monitors and responds to the cell's environment (e.g., to light or to osmotic conditions) via a phospholipid-based signaling pathway. We previously reported that *C. reinhardtii* *bbs* mutants are nonphototactic. Phototaxis is dependent on membrane depolarization and Ca^{2+} influx (Witman, 1993), both of which activate PLD in *C. reinhardtii* (Munnik et al., 2000). Changes in membrane lipid composition could affect the functioning of membrane channels and hence phototaxis. Consistent with this, we previously observed that *bbs4-1* cells with newly formed cilia were able to phototax for over an hour, but lost that ability as the cilia aged and presumably accumulated PLD and acquired an altered membrane lipid composition (Lehtreck et al., 2009).

C. reinhardtii PLD has homologues in mammals and many other ciliated eukaryotes

Our finding that *C. reinhardtii* PLD is apparently a bona fide ciliary protein that cycles through the cilium raises the question of whether a homologue occurs in the cilia of other organisms. Close homologues of *C. reinhardtii* PLD are encoded in the genomes of many vertebrates, nonvertebrate metazoans (with *C. elegans* being a notable exception), and some ciliated protists; they are absent from *Arabidopsis thaliana*, yeasts, and other nonciliated organisms (Fig. S4, A and B). Such a phylogenetic distribution is often indicative of a centriolar or ciliary function of the gene product. The mammalian homologue of PLD is PLD6 (*C. reinhardtii* to *H. sapiens* BLAST E = 8×10^{-37} ; 40% identity); in a phylogenetic analysis PLD6 groups more closely with *C. reinhardtii* PLD than with other murine PLDs (Fig. S4 B). In mice, overexpressed PLD6 localizes to mitochondria and is anchored to the mitochondrial surface by a transmembrane domain (Huang et al., 2011). This transmembrane domain is not predicted in *C. reinhardtii* PLD but, like *C. reinhardtii* AMPK and CAH6, the protein is predicted to be myristoylated and palmitoylated (Fig. S4 C), likely accounting for its association with the ciliary membrane. PLD6 is widely expressed in human tissues; in mice, expression is especially high in lung and testis, tissues that are rich in cilia, and *Pld6*^{-/-} mice are infertile (Huang et al., 2011; Watanabe et al., 2011). It will be of interest to determine if mammalian PLD6 functions also in cilia, and whether BBSome deficiency promotes ciliary accumulation of PLD6. It may be relevant that cilia of *Bbs*^{-/-} mice accumulate unusual intraciliary vesicles at the ciliary tip (Shah et al., 2008); this might be brought about by an excess of ciliary DAG caused by ciliary accumulation of PLD6. Phospholipid signaling also has been implicated in the functioning of olfactory cilia (Klasen et al., 2010), and malfunction of olfactory cilia has been reported in *Bbs8*^{-/-} mice (Tadenev et al., 2011).

BBSome deficiency results in secondary defects in ciliary composition

Quantitative analysis of age-sorted cilia revealed that the ciliary protein CAH6 is progressively lost from *bbs4-1* cilia. Because CAH6 is initially present in mutant cilia, it is unlikely that BBSome defects inhibit CAH6 ciliary import. Rather, the progressive loss of CAH6 from *bbs4-1* cilia is likely to be caused indirectly by BBSome deficiency. Loss of a peripheral membrane protein such as CAH6 could follow from the severe changes in membrane lipid content that occur in *bbs4-1* cilia, presumably as a result of increased ciliary PLD activity; the ability of proteins, such as CAH6, which is predicted to be fatty acid modified (Fig. S4 C), to associate with membranes is influenced by the lipid composition of the membrane (Levental et al., 2010). An altered lipid composition of *bbs* cilia also could contribute in other ways to changes in the pattern of ciliary membrane proteins. Loss of certain GPCRs from cilia has been reported in murine *Bbs*^{-/-} mutants (Berbari et al., 2008b). In light of our data on CAH6 in *bbs* mutant cilia of *C. reinhardtii*, it seems possible that absence of GPCRs from mammalian cilia also could be an indirect consequence of the loss of the BBSome.

BBS likely involves progressive deterioration of ciliary function

The observation that ciliary defects resulting from BBSome deficiency increase with time potentially has implications for the pathomechanism of BBS in mammals. Several features of BBS in murine *Bbs*^{-/-} models or human patients with BBS also increase with time or manifest themselves only years or decades after birth (Beales, 2005). The outer segments of photoreceptor cells in the eye, for example, initially develop and function normally in *Bbs2* and *Bbs4* knockout mice. About 6 wk postnatally, rhodopsin mislocalization becomes apparent and retinal degeneration begins (Nishimura et al., 2004). Therefore, BBSome defects appear to initiate a cascade of events that increasingly deteriorate ciliary protein composition and function. The time scales in which ciliary defects develop in different organisms and different tissues within an organism may be different. Nevertheless, the mechanism by which cells use the BBSome to control ciliary protein transport and to maintain the balance of ciliary signaling proteins might very well be conserved.

Materials and methods

Strains and culture conditions

Wild-type and mutant strains of *C. reinhardtii* were mostly maintained as described in Lechtreck et al. (2009). The *bbs8-1* strain was generated as described in the legend of Fig. S1. The retrograde IFT mutants *fla15* and *fla17-1* were aerated without CO₂ supplement and maintained at 22°C.

Antibodies and Western blotting

Using primers 5'-CGCGAATTCATGGGTTGCGCCAGCTCC-3' and 5'-CGCGAATTCCTAGATGGCAAACCTATGGTGCATG-3', a partial cDNA encoding *C. reinhardtii* PLD was obtained by PCR and cloned into bacterial expression vectors pGEX and pMAL. Truncated PLD fused to maltose-binding protein was used as an antigen to immunize two rabbits, and a fusion protein of truncated PLD and glutathione S-transferase was used to affinity purify the antisera produced by GenScript. Anti-PLD was diluted 1:2,000 for Western blotting and 1:100 for immunofluorescence microscopy. Polyclonal antibodies to *C. reinhardtii* CAH6 (1:2,000) and FAP12 (1:1,000) were provided by J.V. Moroney (Louisiana State University, Baton Rouge, LA) and E. Betleja and D.G. Cole (University of Idaho, Moscow, ID), respectively. Other rabbit polyclonal antibodies used in this study were anti-PKD2 L1 (1:1,000; Huang et al., 2007), anti-IFT140 (1:400), anti-IFT74₂ (1:4,000; Qin et al., 2004), anti-BBS4 (1:800; Lechtreck et al., 2009), anti-DHC1b (1:2,000; Pazour et al., 1999), and anti-D1bLIC (1:1,000; Hou et al., 2004). Mouse monoclonal antibodies used included anti-IC2 (1:100; King and Witman, 1990), anti-IFT139 (1:50), anti-IFT81 (1:250), anti-IFT57 (1:100), and anti-IFT172 (1:50; all Cole et al., 1998); anti-FMG-1 No. 61 (1:1,000; Bloodgood and Salomonsky, 1994); and anti-mastigoneme No. 37 (1:100; Nakamura et al., 1996). Western blots were developed using anti-mouse or anti-rabbit IgG conjugated to horseradish peroxidase (Molecular Probes) and chemiluminescence. An Alpha Innotech FluorChem Q chemiluminescence imaging system was used for documentation and quantification of signals, except for those Western blots including *fla15*, *fla17-1*, and *fla10-1*, which were documented using x-ray film and signals quantified using ImageJ (National Institutes of Health).

Mating experiments and fluorescence microscopy

To generate gametes, cells were grown for 10 d on Tris-acetate-phosphate medium plates (Gorman and Levine, 1965) and resuspended in modified medium I without nitrogen (8 ml/plate; M-N; <http://www.chlamy.org/SG.html>), and the suspension was incubated overnight in constant light with gentle agitation. The next morning, cells were incubated in 20% M-N supplemented with 10 mM Hepes for 3 h. To obtain a series of zygotes of distinct age, gametes were mixed at various consecutive time points. Next, the cells were harvested (3 min at 2,000 rpm, at room temperature); resuspended in 10 mM Hepes, 5 mM MgSO₄, 5 mM EGTA, and 25 mM KCl (HMEK); mixed with an equal volume of 6% formaldehyde/0.1% Nonidet

P-40 in HMEK; and allowed to settle onto poly-L-lysine-coated multi-well slides for ~3–6 min. Slides were then submerged into -20°C methanol for 6 min and air dried. All subsequent steps were performed as described in the following paragraph.

For immunofluorescence microscopy of vegetative cells, cells in HMEK were mixed with an equal volume of 6% formaldehyde/0.5% Nonidet P-40 in HMEK, allowed to settle onto coated slides for 8–15 min, washed with PBS, and air dried. The following primary antibodies and dilutions were used: anti-PLD (1:100), anti-IFT139 (1:1), and anti-acetylated α -tubulin (1:2,000, mouse; Sigma-Aldrich). Specimens were incubated overnight at 4°C with primary antibodies in PBS/2% BSA/0.05% Tween20. Secondary antibodies linked to Alexa Fluor 488, 568, or 594 (1:1,000; Invitrogen) were applied for 90–120 min at room temperature. Specimens were mounted with ProlongGold (Invitrogen). Images were acquired using Axiovision software (Carl Zeiss) and an AxioCam MRm camera (Carl Zeiss) on a microscope (Axioskop 2 plus; Carl Zeiss) equipped with a 100 $\times$ NA 1.4 oil DIC Plan-Apochromat objective (Carl Zeiss) and epifluorescence. Image brightness and contrast were adjusted using Photoshop (CS5.1; Adobe), and figures were assembled using Illustrator (CS5.1; Adobe). Capture times and adjustments were similar for images mounted together. An Axio Imager.Z1 equipped with a 100 $\times$ NA 1.3 EC Plan-Neofluar objective, an Apotome, an axioCam MRm, and Axiovision software (all from Carl Zeiss) was used to obtain optical sections of the immunostained cell shown in Fig. 1 G. Figures were prepared using Adobe Photoshop and Illustrator.

Isolation of cilia

Flagellar isolation using the dibucaine method and flagellar amputation by pH shock were performed as previously described by Witman (1986) and Lefebvre (1995), respectively. In brief, cells were concentrated and repeatedly washed in 10 mM Hepes, pH 7.4, resuspended in HMS (10 mM Hepes, pH 7.4, 5 mM MgSO₄, and 4% wt/vol sucrose), and placed on ice; all subsequent steps were performed at 4°C. 2 ml of dibucaine (25 mM in H₂O) were added per 10 ml of sample; cells were deciliated by rapidly drawing them up and down in a pipette. 20 ml of 0.7 mM EGTA in HMS was then added to the suspension. Cell bodies were removed by differential centrifugation (3 min at 1,150 g); the supernatant was collected and the remaining cell bodies were removed by centrifugation (10 min at 1,700 g) through a sucrose cushion (10 ml 25% sucrose in HSM). The supernatant was harvested and cilia were pelleted at 27,000 g for 20 min. For deciliation by pH shock, 0.5 M acetic acid was added to cells in M medium with rapid stirring until the pH reached 4.2; after 50 s the solution was neutralized using 0.25 N KOH. Cells were immediately placed into fresh culture medium and allowed to regenerate cilia in bright light at RT with shaking (60 rpm). Isolated flagella were extracted with 1% NP-40 in HMDEKP (30 mM Hepes, 5 mM MgSO₄, 1 mM DTT, 0.5 mM EGTA, 25 mM KCl, and 1% protease inhibitor cocktail [Sigma-Aldrich]) for 20 min on ice, and axonemes and membrane-plus-matrix were separated by centrifugation (27,000 g for 10 min). Alternatively, cilia were extracted with 1% Triton X-114, which allows subsequent separation of the membrane and matrix fractions by phase partitioning. After removing the axonemes, the supernatant was incubated briefly at 37°C and phases were separated by centrifugation (3,300 g for 10 min at RT). The aqueous phase was treated with 1% Triton X-114, the detergent phase was diluted with HMDEKP, and the phase separation was repeated to yield the final aqueous and detergent phases.

For temperature-shift experiments with *fla10-1*, cells were concentrated and incubated in Modified Medium I (Witman, 1986) with gentle shaking (60 rpm) either at room temperature (~22°C) or at 32°C in an incubator. For time course experiments, equal aliquots of concentrated *fla10-1* cells were transferred at various time points from 22 to 32°C and incubated for the time period indicated. Then, cilia from all samples were isolated simultaneously using warm (32°C) 10 mM Hepes to wash cells maintained at the restrictive temperature. All experiments using the *dhc1b-2* and other IFT mutants were performed at or below 22°C.

Lipidomic and metabolomic analyses of cilia

We used two distinct approaches for the lipidomic analysis of cilia, liquid chromatography coupled to mass spectrometry and an approach based solely on mass spectrometry. In brief, cilia were extracted with methanol/chloroform and lipid extracts were separated by gradient elution normal phase HPLC and the analytes detected by positive atmospheric pressure chemical ionization. For metabolomic analysis by gas chromatography coupled to mass spectrometry, samples were derivatized to their trimethylsilyl derivatives. Chromatography was on a 30-M DB-5MS column. Full scan

electron ionization mass spectrometry was performed using a Quattro II mass spectrometer (Waters). The accuracy of quantitation is $\pm 10\%$.

For quantitative analysis of phospholipids, $\sim 100 \mu\text{g}$ of cilia (wet weight) from each sample was subjected to lipid extraction. In brief, the sample was dissolved in $200 \mu\text{l}$ of liquid chromatography grade water. Samples were spiked with $10 \mu\text{l}$ of a mixture of internal standards containing $20 \text{ pmol DAG } 17:0-17:0$, $24 \text{ pmol PA } 17:0-17:0$, $52 \text{ pmol PE } 17:0-17:0$, $7.5 \text{ pmol phosphatidylglycerol (PG) } 17:0-17:0$, $43 \text{ pmol phosphatidylserine (PS) } 17:0-17:0$, $40 \text{ pmol phosphatidylcholine (PC) } 18:3-18:3$, $54 \text{ pmol phosphatidylinositol (PI) } 17:0-17:0$, $20 \text{ pmol ceramide (Cer) } 18:1;2/17:0;0$, $40 \text{ pmol sphingomyelin (SM) } 18:1;2/17:0$, $20 \text{ pmol galactosylceramide (GalCer) } 18:1;2/12:0$, and $20 \text{ pmol lactosylceramide (LacCer) } 18:1;2/12:0$. $265 \mu\text{l}$ of methanol was added to the mixture and agitated for 10 min for homogenization. Then, $730 \mu\text{l}$ of chloroform were added and the mixture was agitated for an additional 1 h. The lower organic phase was collected and evaporated in a Speedvac concentrator at room temperature. Lipid extracts were dissolved in $100 \mu\text{l}$ of 0.1% methylamine in methanol and subjected to quantitative lipid analysis on a hybrid QSTAR Pulsar i quadrupole time-of-flight mass spectrometer (MDS Sciex). Samples were infused with a TriVersa NanoMate robotic nanoflow ion source (Advion Biosciences, Inc.). DAG, PA, PS, PE, PI, and PG species were quantified by negative ion mode multiple precursor ion scanning analysis (Ejsing et al., 2006).

Miscellaneous

For mass spectrometric identification of CAH6, bands of interest were excised from silver-stained gels and digested overnight with proteomics-grade trypsin, and the eluted peptides were analyzed using a matrix-assisted laser desorption/ionization time-of-flight mass spectrometer (Kratos Axima QIT; Shimadzu Instruments). Peptides were analyzed in positive ion mode in mid-mass range ($700-3,000 \text{ D}$; Lechtreck et al., 2009).

CLUSTALW at the default settings was used to align multiple PLD sequences and for tree construction.

Online supplemental material

Fig. S1 A shows a molecular map of *bbs8-1*; the sequence of CAH6 is depicted in Fig. S1 B. Fig. S2 analyzes ciliary length and IFT in *dhc1b-2* and the *bbs* mutant strains and shows protein content of wild-type and *bbs* mutant cilia and cell bodies. Fig. S3 shows ciliary numbers and length for *fla10-1* and *dhc1b-2* and compares ciliary regeneration in wild type and *bbs4-1*. An alignment and a phylogenetic analysis of various PLDs is shown in Fig. S4, A and B, respectively. Fig. S4 C shows the predicted fatty acid modification sites for PLD, AMPK, and CAH6. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201207139/DC1>.

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