

2012-10-01

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Engel BD, Ishikawa H, Wemmer KA, Geimer S, Wakabayashi K, Hirono M, Craige B, Pazour GJ, Witman GB, Kamiya R, Marshall WF. (2012). The role of retrograde intraflagellar transport in flagellar assembly, maintenance, and function. *Cell and Developmental Biology Publications*. <https://doi.org/10.1083/jcb.201206068>. Retrieved from https://escholarship.umassmed.edu/cellbiology_pp/125

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The role of retrograde intraflagellar transport in flagellar assembly, maintenance, and function

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The maintenance of flagellar length is believed to require both anterograde and retrograde intraflagellar transport (IFT). However, it is difficult to uncouple the functions of retrograde transport from anterograde, as null mutants in dynein heavy chain 1b (DHC1b) have stumpy flagella, demonstrating solely that retrograde IFT is required for flagellar assembly. We isolated a *Chlamydomonas reinhardtii* mutant (*dhc1b-3*) with a temperature-sensitive defect in DHC1b, enabling inducible inhibition of retrograde IFT in full-length flagella. Although *dhc1b-3* flagella at the nonpermissive temperature (34°C)

showed a dramatic reduction of retrograde IFT, they remained nearly full-length for many hours. However, *dhc1b-3* cells at 34°C had strong defects in flagellar assembly after cell division or pH shock. Furthermore, *dhc1b-3* cells displayed altered phototaxis and flagellar beat. Thus, robust retrograde IFT is required for flagellar assembly and function but is dispensable for the maintenance of flagellar length. Proteomic analysis of *dhc1b-3* flagella revealed distinct classes of proteins that change in abundance when retrograde IFT is inhibited.

Introduction

Cilia and flagella are dynamic structures whose assembly and maintenance require an active transport process known as intraflagellar transport (IFT; Kozminski et al., 1993; Pedersen and Rosenbaum, 2008; Scholey, 2008; Ishikawa and Marshall, 2011). Intraflagellar transport is mediated by ~20 IFT proteins (Piperno and Mead, 1997; Cole et al., 1998), which assemble into macromolecular complexes called IFT particles. The IFT particles then associate into linear arrays known as IFT trains (Pigino et al., 2009), which move processively from the base of the flagellum out to the tip. This anterograde transport is driven by kinesin-2, a heterotrimeric complex composed of the FLA10 and FLA8 motor subunits (Walther et al., 1994) and the kinesin-associated protein KAP (Cole et al., 1993; Mueller et al., 2005). After their anterograde motion to the flagellar tip, IFT particles rearrange into a new set of IFT trains that move back to the base

of the flagellum. This retrograde transport is powered by cytoplasmic dynein 1b, a large complex composed of the heavy chain motor subunit DHC1b (Pazour et al., 1999a; Porter et al., 1999; Signor et al., 1999) and numerous smaller components including D1bLIC (Perrone et al., 2003; Schafer et al., 2003; Hou et al., 2004; Hao et al., 2011a), FAP133 (Rompolas et al., 2007), and LC8 (Pazour et al., 1998). Both the precursors for flagellar assembly and the breakdown products of flagellar turnover are thought to associate with IFT particles (Qin et al., 2004; Hao et al., 2011b).

The primary evidence that anterograde IFT supplies precursors for flagellar assembly to the growing tip of the flagellum comes from analysis of mutations in the anterograde motor kinesin-2. Using a *fla10* temperature-sensitive (*ts*) allele, it was shown that kinesin activity is required for IFT train motion (Kozminski et al., 1995), flagellar assembly and length maintenance (Kozminski et al., 1995), and the incorporation of new tubulin into the flagellar tip during steady-state turnover

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Abbreviations used in this paper: 2D-DIGE, two-dimensional difference in gel electrophoresis; CDS, coding sequence; DIC, differential interference contrast; IFT, intraflagellar transport; KAP, kinesin-associated protein; NaPPi, sodium pyrophosphate; TAP, Tris-acetate-phosphate; TEM, transmission electron microscopy; TIRF, total internal reflection fluorescence; *ts*, temperature-sensitive; UTR, untranslated region.

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(Marshall and Rosenbaum, 2001). Furthermore, changes in protein abundance in *fla10* flagella were used for the initial identification of the IFT proteins (Piperno and Mead, 1997; Cole et al., 1998). Thus, our understanding of anterograde IFT has resulted in large part from the availability of a conditional mutation in the anterograde motor that allows inducible shutoff of motor function.

Compared with anterograde transport, our understanding of the functional role of retrograde transport is less well developed. *Chlamydomonas* mutant strains with null deletions in DHC1b (*dhc1b-1*, Pazour et al., 1999a; *stf-1*, *stf-2*, Porter et al., 1999) have stumpy flagella, swollen with IFT and axonemal proteins that have apparently been stranded at the flagellar tip in the absence of return transport. Based on this phenotype, it was concluded that an important function of retrograde transport is to carry back IFT particles and kinesin motors so they can execute a new cycle of anterograde transport. It is believed that flagellar length cannot be maintained without this recycling, as the anterograde transport machinery becomes sequestered in the flagellum.

Unfortunately, constitutive mutations in DHC1b only reveal endpoint phenotypes and do not provide information about the dynamics by which these phenotypes arise. Consequently, it is impossible to determine if the swollen stumpy flagella of *dhc1b-1* reflect a role of dynein in the initial assembly of flagella or in their subsequent maintenance. In the case of anterograde transport, the *ts fla10* mutant demonstrated the requirement of anterograde IFT for flagellar maintenance by dynamically shortening its flagella when the mutant was shifted to the nonpermissive temperature (Kozminski et al., 1995). However, the only *ts* dynein mutant that has been isolated up to this point (*dhc1b-2*, Pazour et al., 1999b; Witman, 2011) does not clearly reveal the role of retrograde IFT in flagellar maintenance, as even at the permissive temperature it has impaired dynein function and short flagella full of accumulated IFT material (Fig. S1).

Here, we describe a new conditional mutation in dynein, *dhc1b-3*, which enables the inducible decoupling of retrograde IFT from anterograde transport. Strains carrying this mutation have full-length flagella and nearly wild-type levels of IFT at the permissive temperature, but lose the majority of retrograde IFT events after a shift to the nonpermissive temperature. Anterograde IFT also becomes partially decreased upon temperature-shifting the *dhc1b-3* mutant; however, flagellar length is maintained with only mild shortening for many hours until cell division, when new flagella are unable to assemble. This result is dramatically distinct from the immediate flagellar shortening seen in *fla10*, and suggests that robust retrograde IFT is necessary for flagellar assembly but not length maintenance. We further analyzed the role of retrograde IFT by comparative proteomics, which revealed complex patterns of protein accumulation and depletion in *dhc1b-3* flagella. In addition, *dhc1b-3* cells at the nonpermissive temperature displayed changes in phototaxis, swimming speed, and flagellar beat frequency. Taken together, our results indicate that retrograde IFT is not simply a means to recycle anterograde IFT trains, but plays a multifaceted role in maintaining flagellar composition and function.

Results

Isolation and cloning of *dhc1b-3*, a temperature-sensitive DHC1b mutation

In an effort to obtain conditional mutations affecting retrograde IFT, we performed UV mutagenesis of *Chlamydomonas* cells, followed by a phenotypic screen for *ts* defects in cell motility (Fig. S2 E). Of the 122 punctate colonies isolated for further screening, 68 were bald, 13 had jerky swimming, 6 were paralyzed, 17 were sick, and 18 swam normally (screen false hits). Only 3 isolates exhibited *ts* motility defects. Two of these strains had conditional flagellar paralysis and were not further studied, whereas a single strain showed a robust and reversible defect in flagellar assembly. At 21°C, this mutant had full-length flagella containing a normal distribution of IFT proteins (Fig. 1 A) and appeared to have no ultrastructural defects in the axoneme or basal body (Fig. 1 D). However, after incubation for a day at 34°C, IFT proteins accumulated in flagella (Fig. 1, B and E) and flagella became stumpy and swollen with IFT material (Fig. 1, C and F). These stumpy flagella persisted until cells were shifted back to 21°C, at which temperature flagella regrew after several hours. The accumulation of IFT proteins in the mutant's flagella at the nonpermissive temperature was typical of a retrograde IFT defect and was highly reminiscent of null mutants in DHC1b, the heavy chain of the retrograde dynein motor (*dhc1b-1*, Pazour et al., 1999a; *stf-1*, *stf-2*, Porter et al., 1999), which have stumpy swollen flagella.

During initial analysis of tetrads obtained from backcrosses (Harris et al., 2009), we found that the mutant phenotype was tightly linked to the mating-type locus on chromosome 6. To perform higher resolution mapping, we crossed the mutant to the S1C5 polymorphic strain (Gross et al., 1988) and used PCR-based SNP analysis on the tetrad progeny to check for recombination between the mutation and several genetic markers (Fig. S2 A). This analysis tracked the mutation to a small region at the end of chromosome 6. Among the four potential flagellar genes in this region was DHC1b, our leading candidate based on phenotypic similarities to the null *dhc1b-1* mutant.

Sequencing the entire length of the DHC1b cDNA from the mutant strain revealed a single base-pair mutation at 12313 bp in the coding sequence (Fig. S3 A). This A-to-T substitution results in a single amino acid switch from isoleucine to phenylalanine at position 4105 of the amino acid sequence (Fig. S3 B). The mutation occurs in the "C-sequence," the C-terminal domain beyond the six ATPase domains (Kon et al., 2011, 2012). The C-sequence is present in the dynein heavy chains of many species (Numata et al., 2011), but is absent from budding yeast (Carter et al., 2011; Schmidt et al., 2012). ClustalX alignment (Larkin et al., 2007) of DHC1b C-sequences from several organisms revealed that the mutation occurs in a fully conserved hydrophobic residue in the middle of a small conserved hydrophobic motif (Fig. S3 C). Thus, the replacement of Ile with the bulkier Phe residue could presumably impede protein stability, a defect that would be exacerbated by the temperature shift. It is also possible that the mutation affects motor function, as the C-sequence is crucial for dynein processivity (Numata et al., 2011).

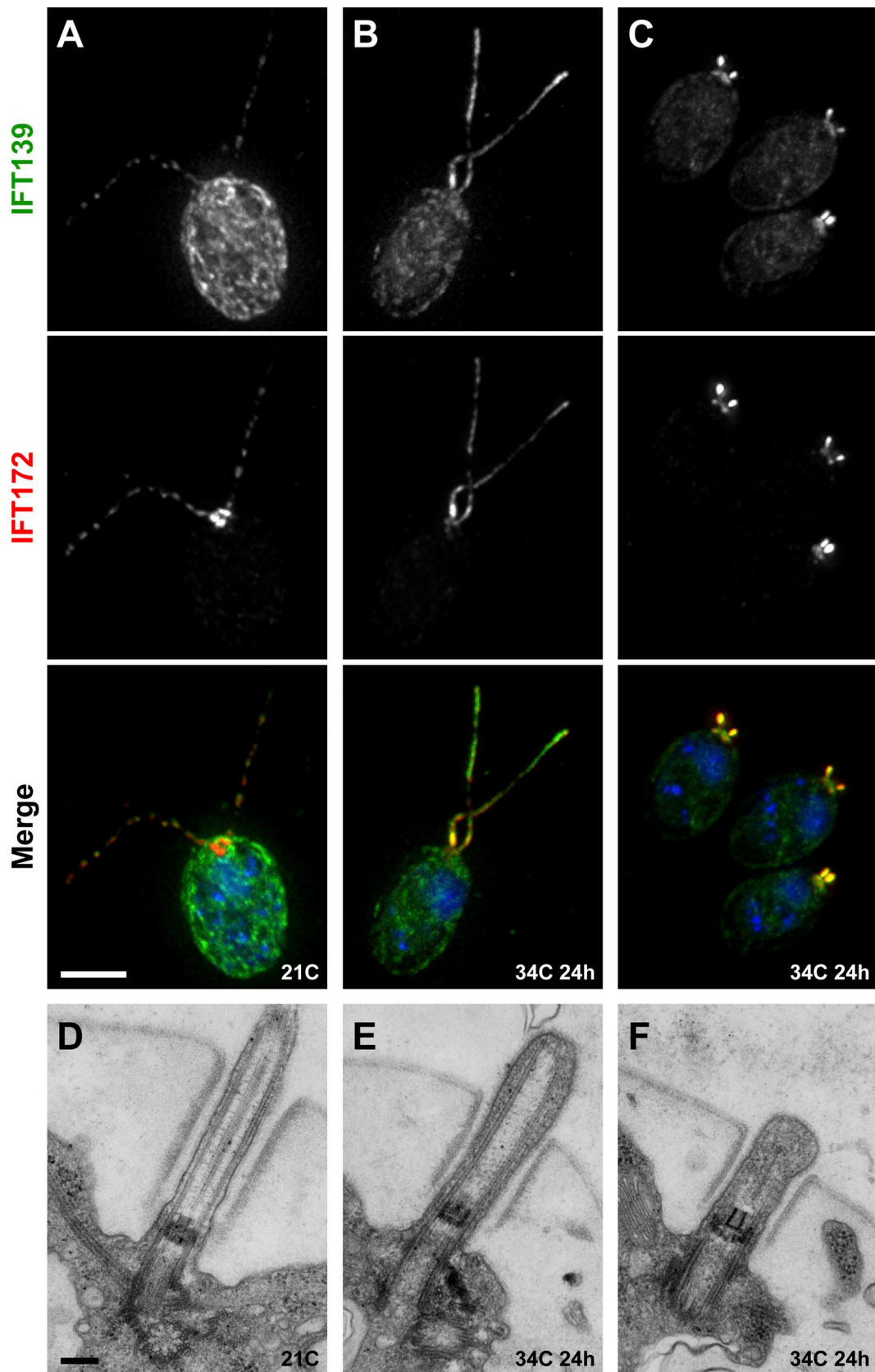


Figure 1. **Temperature-sensitive “retrograde IFT” phenotype of isolated mutant strain *dhc1b-3*.** (A–C) Immunofluorescence and (D–F) transmission electron microscopy (TEM) of the *ts* flagellar assembly mutant. Green, IFT139; red, IFT172; blue, DNA. At 21°C, the mutant appeared to have a regular distribution of IFT proteins along the flagella (A) and showed no discernible defects in axoneme or basal body ultrastructure (D). After 24 h at 34°C, flagella were either long (B) or stumpy (C) and in both cases showed a strong accumulation of IFT proteins. This accumulation of IFT proteins at 34°C was also apparent by TEM (E and F) as a buildup of electron-dense material between the axoneme and the flagellar membrane. Bars: (A–C) 5 μ m; (D–F) 200 nm.

We named the mutant strain *dhc1b-3* and developed a PCR assay to easily track the mutation, verifying that there is heterogeneity between *dhc1b-3* and wild-type sequences at the mutated site (Fig. S2, B and C). To confirm that the *dhc1b-3* phenotype is due to an impairment of the DHC1b gene, we created diploid strains by mating *dhc1b-3* to the *dhc1b-1* null mutant (Fig. S2 D). *dhc1b-1* is a recessive mutation and would therefore show no phenotype if wild-type DHC1b function was present. The *dhc1b-1 dhc1b-3* diploids exhibited conditional loss of flagella at 34°C with similar kinetics to the parental *dhc1b-3* strain. Thus, the *dhc1b-3* and *dhc1b-1* strains do not complement, indicating that the *dhc1b-3* phenotype is caused by a specific effect on the DHC1b gene.

Retrograde IFT is reduced in *dhc1b-3* flagella

To determine how the *dhc1b-3* mutation affects IFT, we imaged the flagella of live cells by two different light microscopy techniques. After crossing *dhc1b-3* cells to a strain expressing the KAP subunit tagged with green fluorescent protein (KAP-GFP; Mueller et al., 2005), we visualized KAP-GFP movement using total internal reflection fluorescence (TIRF) microscopy (Fig. 2 A and Videos 1–3; for review of technique see Engel et al., 2009a). One distinct feature of KAP-GFP is that there is a high level of background fluorescence within flagella instead of clear retrograde traces (Engel et al., 2009b). Although *dhc1b-3* KAP-GFP looked similar to wild-type at 21°C, anterograde IFT became less frequent at 34°C and the background fluorescence decreased. Interestingly, in the absence of strong background signal, we observed some KAP-GFP proteins undergoing bidirectional oscillatory movements, a behavior we have never witnessed in wild-type KAP-GFP flagella. To better understand how retrograde transport is affected in *dhc1b-3* cells, we also visualized IFT using differential interference contrast (DIC) microscopy (Fig. 2 B and Videos 4–6; for reviews of technique see Kozminski, 1995; Dentler et al., 2009). At 21°C, *dhc1b-3* IFT appeared similar to wild-type, though retrograde traces were slightly less processive. However, after 6–12 h at 34°C, retrograde transport events were dramatically slower and less frequent. Mirroring the KAP-GFP imaging, anterograde IFT frequency also decreased at 34°C.

After imaging, we quantified IFT speeds and frequencies by kymograph analysis (Fig. 2, C and D). Incubating wild-type cells at 34°C for 6 or 12 h had a negligible effect on IFT. Anterograde train speed was 2.5 μm/s, retrograde speed was 4.4 μm/s, anterograde frequency was 1.6 trains/s (1.1 trains/s for TIRF), and retrograde frequency was 2.3 trains/s. When *dhc1b-3* cells were grown at 21°C, anterograde speed and frequency were unchanged from wild-type levels, but retrograde speed was decreased to 3.3 μm/s and retrograde frequency dropped to 1.7 trains/s. Retrograde train speed in *dhc1b-3* flagella was further reduced to 2.1 μm/s after 6 h at 34°C and 1.7 μm/s after 12 h. Most dramatically, *dhc1b-3* retrograde frequency dropped to 0.8 trains/s and 0.5 trains/s after 6 and 12 h incubation at 34°C, respectively. The impact of the *dhc1b-3* mutation on anterograde IFT was less severe. Anterograde speed was only marginally affected, decreasing to 2.2 μm/s (1.8 μm/s for TIRF) after

12 h at 34°C. However, anterograde frequency experienced a significant reduction to 1.0 trains/s (0.9 trains/s for TIRF) after 6 h at 34°C and 0.9 trains/s (0.5 trains/s for TIRF) after 12 h. IFT train size (Engel et al., 2009b) was not measured in these experiments due to inadequate GFP expression in the *dhc1b-3* KAP-GFP strain.

Unlike the *ts* kinesin mutant *fla10*, IFT does not completely stop in *dhc1b-3* cells during the first 12 h at 34°C. Instead, retrograde IFT speed and frequency are greatly reduced, coupled with a moderate decrease in anterograde IFT frequency. How this IFT defect may be linked to the *dhc1b-3* flagellar phenotype is explored further in the Discussion.

dhc1b-3 has distinct kinetics of flagellar loss

Although several characterized *Chlamydomonas* mutants have *ts* defects in flagellar assembly (most notably the *fla* mutants; Adams et al., 1982), the kinetics of flagellar loss in *dhc1b-3* are strikingly different. Although *fla10* flagella began shortening immediately at 34°C and were lost within 5 h (Fig. 3 C; Kozminski et al., 1995; Marshall and Rosenbaum, 2001), vegetative *dhc1b-3* cells maintained nearly full-length flagella for over a day at 34°C before finally losing them (Fig. 3 A). Unlike *fla10*, only mild flagellar shortening was observed in *dhc1b-3* cells until after the majority of the population had undergone deflagellation. The difference between *fla10* and *dhc1b-3* shortening kinetics suggests that although both anterograde and retrograde IFT are necessary for flagellar assembly, robust retrograde IFT may not be required for the maintenance of flagellar length.

In synchronized log growth-phase cultures, *dhc1b-3* cells were flagellated at both 21 and 34°C until flagella were resorbed for cell division. After division, flagella regrew at 21°C but were unable to regrow at 34°C (Fig. S4 A). This implies that *dhc1b-3* cells can maintain established flagella at 21 and 34°C, but can only assemble new flagella at 21°C. To further explore this hypothesis, we forced vegetative cells to undergo synchronous differentiation into gametes by nitrogen starvation (Harris et al., 2009). Incubation of vegetative *dhc1b-3* cells for 4–8 h at 34°C before resuspending cells in nitrogen-free media inhibited flagellar assembly in the resulting gametes (Fig. 3 B). This again suggests that after several hours at 34°C, *dhc1b-3* cells cannot assemble new flagella.

We created a *fla10 dhc1b-3* double mutant via mating and found that this strain has identical kinetics of flagellar shortening and loss at 34°C as the *fla10* single mutant (Fig. 3 C). Thus, kinesin inhibition is epistatic to dynein inhibition for the maintenance of flagellar length. This result is not surprising, as kinesin-powered anterograde transport brings the axonemal precursors required for length maintenance to the flagellar tip. One caveat is that the kinetics of kinesin inactivation in *fla10* may be faster than the kinetics of *dhc1b-3* dynein inhibition.

We next observed the effects of the *dhc1b-3* mutation on flagellar assembly more directly by using pH shock regeneration. At 21°C, the extent (Fig. 4 A) and kinetics (Fig. S4 B) of *dhc1b-3* flagellar regeneration were comparable to wild-type. In agreement with the gametogenesis measurements (Fig. 3 B), *dhc1b-3* cells that were incubated at 34°C for 6 h before pH

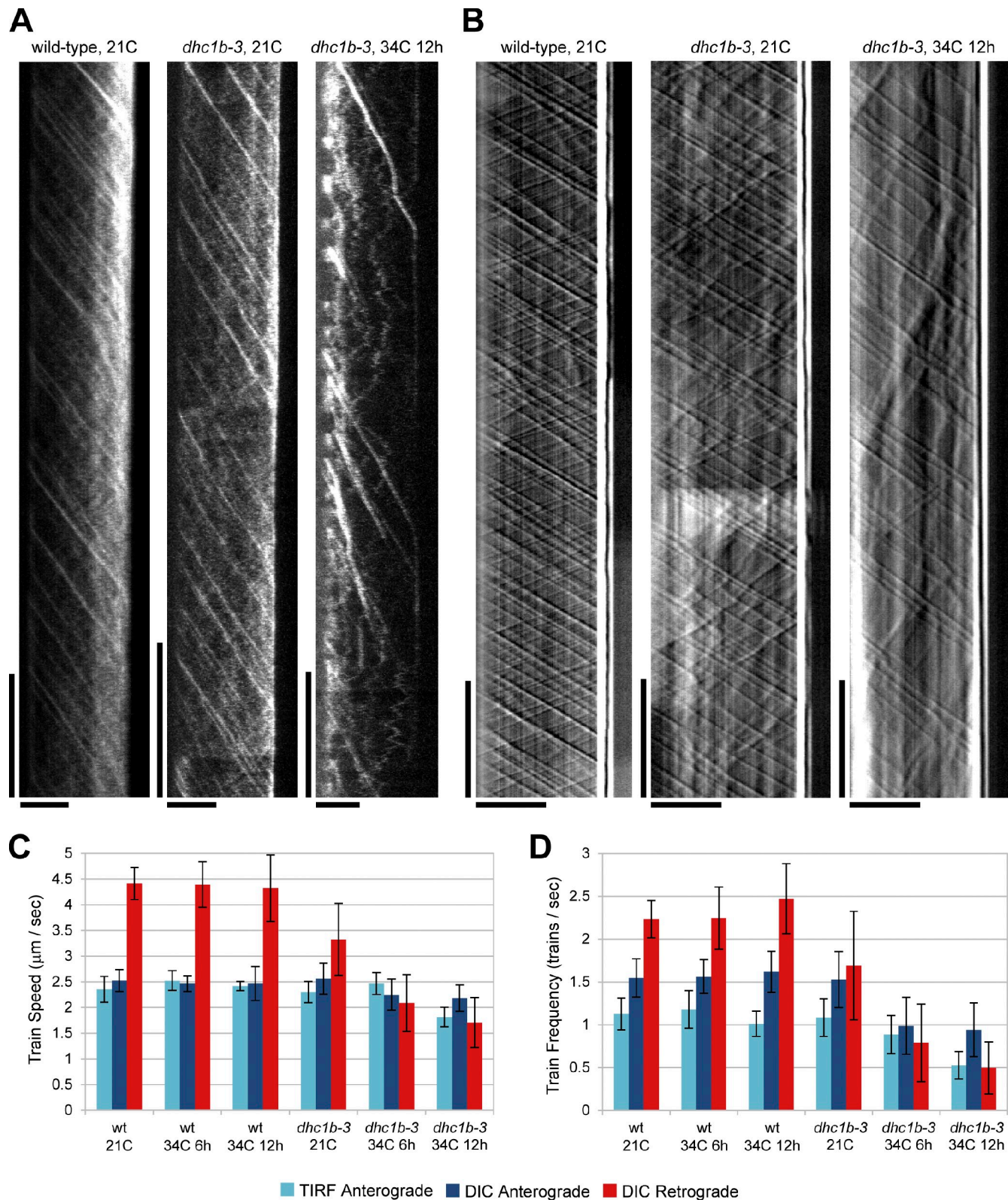


Figure 2. **TIRF and DIC microscopy measurements of IFT in wild-type and *dhc1b-3* cells at 21 and 34°C.** (A) Kymographs generated from TIRF time series of wild-type and *dhc1b-3* cells expressing KAP-GFP at 21 and 34°C, as indicated. (B) Kymographs generated from DIC time series of wild-type and *dhc1b-3* cells at 21 and 34°C. For A and B, the x-axis is distance along the length of the flagellum (the flagellar base is on the left and the flagellar tip is on the right) and the y-axis is time (earliest times are at the top). Bars (vertical) 5 s; (horizontal) 5 μm. Quantification of IFT train speeds (C) and frequencies (D) from the TIRF and DIC datasets. Error bars indicate standard deviation. In *dhc1b-3* flagella, retrograde IFT is moderately reduced at 21°C and strongly inhibited at 34°C. The large drop in *dhc1b-3* retrograde IFT at 34°C is accompanied by a decrease in anterograde IFT frequency. The lower frequency of anterograde trains measured by TIRF vs. DIC likely means that the KAP-GFP fluorescence of some trains was not detected. $N_{\text{flagella}} = 231$ (TIRF), 109 (DIC).

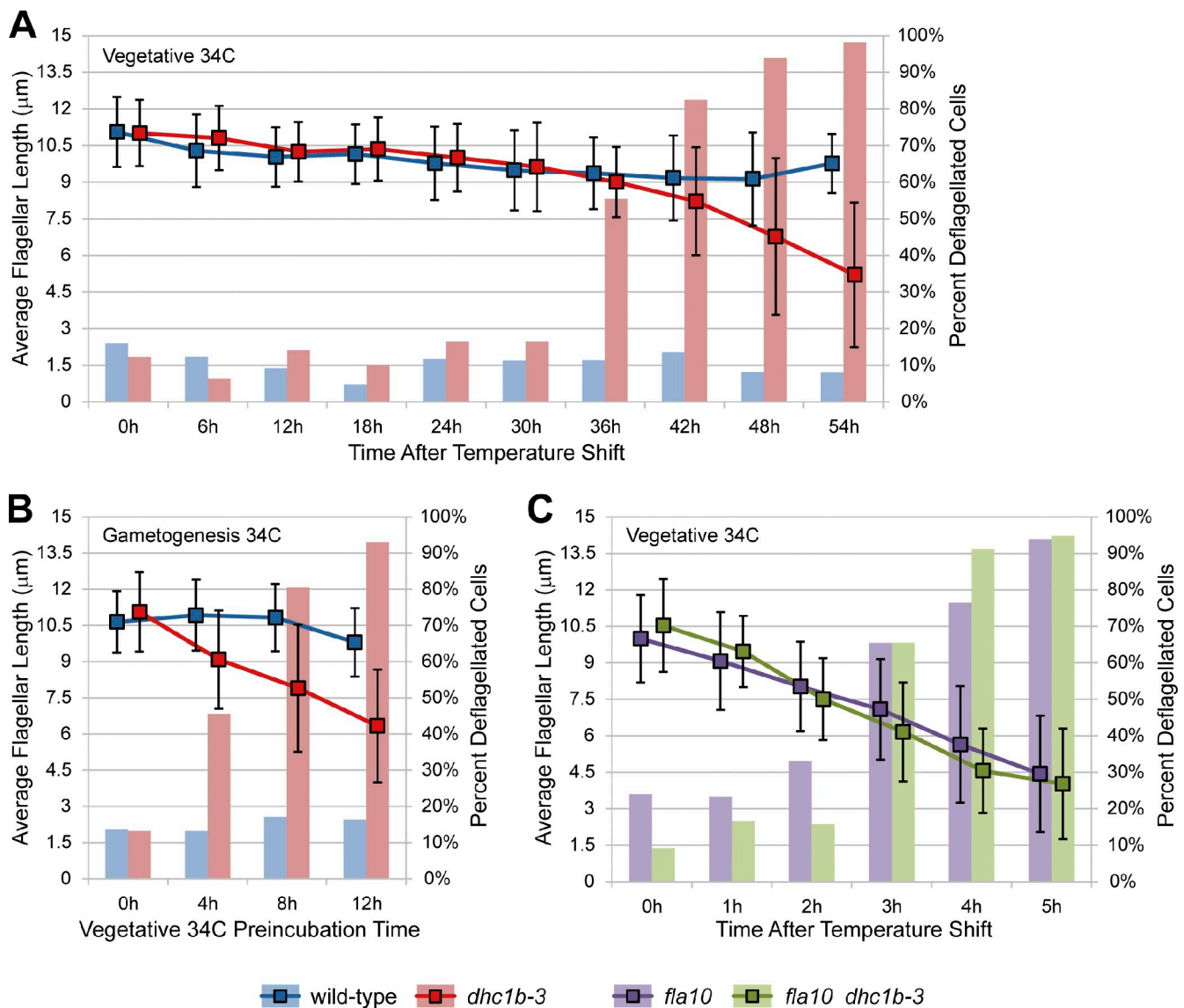


Figure 3. Kinetics of *dhc1b-3* flagellar loss in asynchronous vegetative cells, during gametogenesis, and in *fla10 dhc1b-3* double mutants. Bar graphs depict the percentage of deflagellated cells, and line graphs show the average flagellar length of flagellated cells. Error bars indicate standard deviation. (A) Vegetative wild-type and *dhc1b-3* cells were grown in TAP media for 2 d at 21°C (dense, post-log phase cultures) and then shifted to 34°C. In this experiment, *dhc1b-3* flagella were maintained with only mild shortening (10–20%) for 30 h. In less dense, rapidly dividing cultures, some *dhc1b-3* cells lost flagella as early as 18 h at 34°C (not depicted). The cause of this variability is unknown, but it is likely related to the growth phase of the culture. It was not determined whether flagellar loss in these asynchronous cultures was due to cell division (as shown in Fig. S4 A) or other factors, such as retrograde IFT dropping below a critical threshold. (B) Vegetative cells were grown in TAP media at 21°C for 1 d, incubated for increasing periods of time at 34°C, and then induced to differentiate synchronously into gametes by diluting cells into M-N media followed by 18 h incubation at 34°C. Pre-incubation for 4–8 h at 34°C inhibited the assembly of new flagella in *dhc1b-3* gametes. (C) Vegetative *fla10* and *fla10 dhc1b-3* double mutant strains were grown in TAP media for 2 d at 21°C and then shifted to 34°C. Both strains had the same kinetics of flagellar shortening and loss, which was far more rapid than flagellar loss in the *dhc1b-3* single mutant. $N_{\text{flagella}} = 1,849$ (A), 833 (B), 940 (C) from single experiments.

shock showed a strong flagellar assembly defect during 34°C regeneration, whereas preincubation for 12 h at 34°C completely inhibited regeneration (Fig. 4 A). Furthermore, when *dhc1b-3* cells were incubated at 34°C for increasing periods of time before pH shock, and then allowed to recover at 21°C, flagellar assembly was delayed and proceeded at a slower rate (Fig. S4 B). These results again illustrate the strict requirement for dynein function in flagellar assembly.

Previous studies have indicated a role for IFT in active disassembly of flagella, as the rate of sodium pyrophosphate (NaPPi)-induced flagellar shortening is decreased in *fla10* cells

at the nonpermissive temperature (Pan and Snell, 2005). Because *fla10* abrogates both anterograde and retrograde transport, this result does not reveal which phase of IFT is involved. One appealing hypothesis is that retrograde transport plays a key role in disassembly by facilitating the removal of flagellar proteins. However, we found that the rate and extent of NaPPi-induced flagellar shortening was not reduced in the *dhc1b-3* mutant at 34°C compared with wild-type cells grown at 34°C (Fig. 4 B), indicating that retrograde IFT may not be required for active disassembly. Thus, the increased rate of flagellar shortening in the presence of IFT is likely due to anterograde

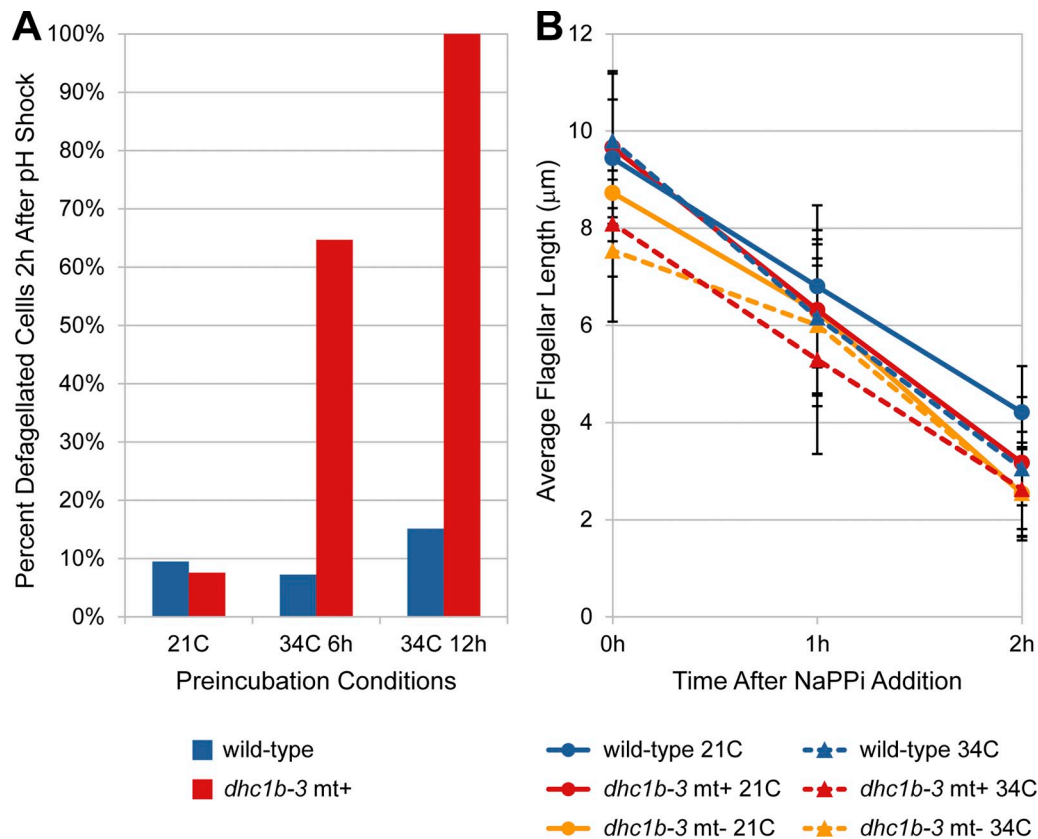


Figure 4. **pH shock-induced flagellar regeneration and NaPPI-induced flagellar shortening in wild-type and *dhc1b-3* cells at 21 and 34°C.** (A) Wild-type and *dhc1b-3* cells were incubated at 21 or 34°C (6 h, 12 h), deflagellated with pH shock, allowed to recover at 21 or 34°C, then scored after 2 h recovery for the percentage of deflagellated cells (regeneration failure). Although the extent of *dhc1b-3* flagellar regeneration at 21°C was similar to wild-type, *dhc1b-3* cells showed strong flagellar assembly defects at 34°C. $N_{\text{cells}} = 2,335$ from two experiments. (B) Cells were grown in M media, incubated for 12 h at 21 or 34°C, and then treated with 20 mM NaPPI. The flagella of wild-type and *dhc1b-3* cells (mt+ and mt- strains) shortened with similar rates ($\sim 3 \mu\text{m}/\text{h}$) at both 21 and 34°C. Thus, NaPPI-induced flagellar shortening may be independent of retrograde IFT. The starting flagellar lengths of both mutant and wild-type strains were slightly shorter than normal, likely due to unknown variations in the culture conditions. Error bars indicate standard deviation. $N_{\text{flagella}} = 540$ from a single experiment.

transport, which has been shown to carry the microtubule-depolymerizing factor kinesin-13 into flagella in response to NaPPI treatment (Piao et al., 2009).

The depletion of dynein from *dhc1b-3* flagella is correlated with the accumulation of IFT proteins, but not kinesin

Given the dramatic reduction in *dhc1b-3* retrograde IFT, we decided to check the flagellar abundance of the retrograde dynein motor and IFT proteins by immunofluorescence microscopy (Fig. 5 A). Relative to wild-type, DHC1b fluorescence was markedly decreased in *dhc1b-3* flagella at both 21°C and after 18 h at 34°C, though some protein remained localized around the basal bodies. This drop in DHC1b abundance was mirrored by the flagellar depletion of D1bLIC, a light intermediate chain of cytoplasmic dynein 1b (Hou et al., 2004). At 34°C, D1bLIC was also displaced from the basal bodies of many *dhc1b-3* cells, localizing to punctate spots throughout the cytoplasm. As dynein fluorescence decreased, the intensities of IFT-B component IFT81 and IFT-A protein IFT139 increased. This result is not unexpected and indeed has been predicted based on the accumulation of IFT proteins in the stumpy flagella of DHC1b-null

mutants (Pazour et al., 1999a; Porter et al., 1999) and the short flagella of *fla14* (deletion of the dynein light chain LC8; Pazour et al., 1998). Nonetheless, to our knowledge this is the first direct demonstration that the induced depletion of cytoplasmic dynein 1b from full-length flagella (and subsequent drop in retrograde IFT, Fig. 2) results in the flagellar accumulation of IFT proteins.

To expand our investigation to a wider scope of proteins and time points, we performed Western blot analysis of isolated flagella from wild-type and *dhc1b-3* cells (Fig. 5 B). DHC1b and D1bLIC levels were greatly reduced in *dhc1b-3* flagella at 21°C, and decreased even further after the mutant was incubated at 34°C. In agreement with the immunofluorescence data, this depletion of dynein was accompanied by an accumulation of IFT-B proteins IFT27, IFT81, and IFT172 (only at 34°C), as well as IFT-A proteins IFT122 and IFT140 (at both 21 and 34°C). We saw no change in BBS4, a component of the BBSome that is trafficked by IFT (Lehtreck et al., 2009). The levels of axonemal proteins α -tubulin and Rib43a (protofilament ribbons; Norrander et al., 2000) remained constant, although ODA-IC2 (intermediate chain of outer arm dynein, ODA6; Mitchell and Kang, 1991) appeared to accumulate in *dhc1b-3* flagella.

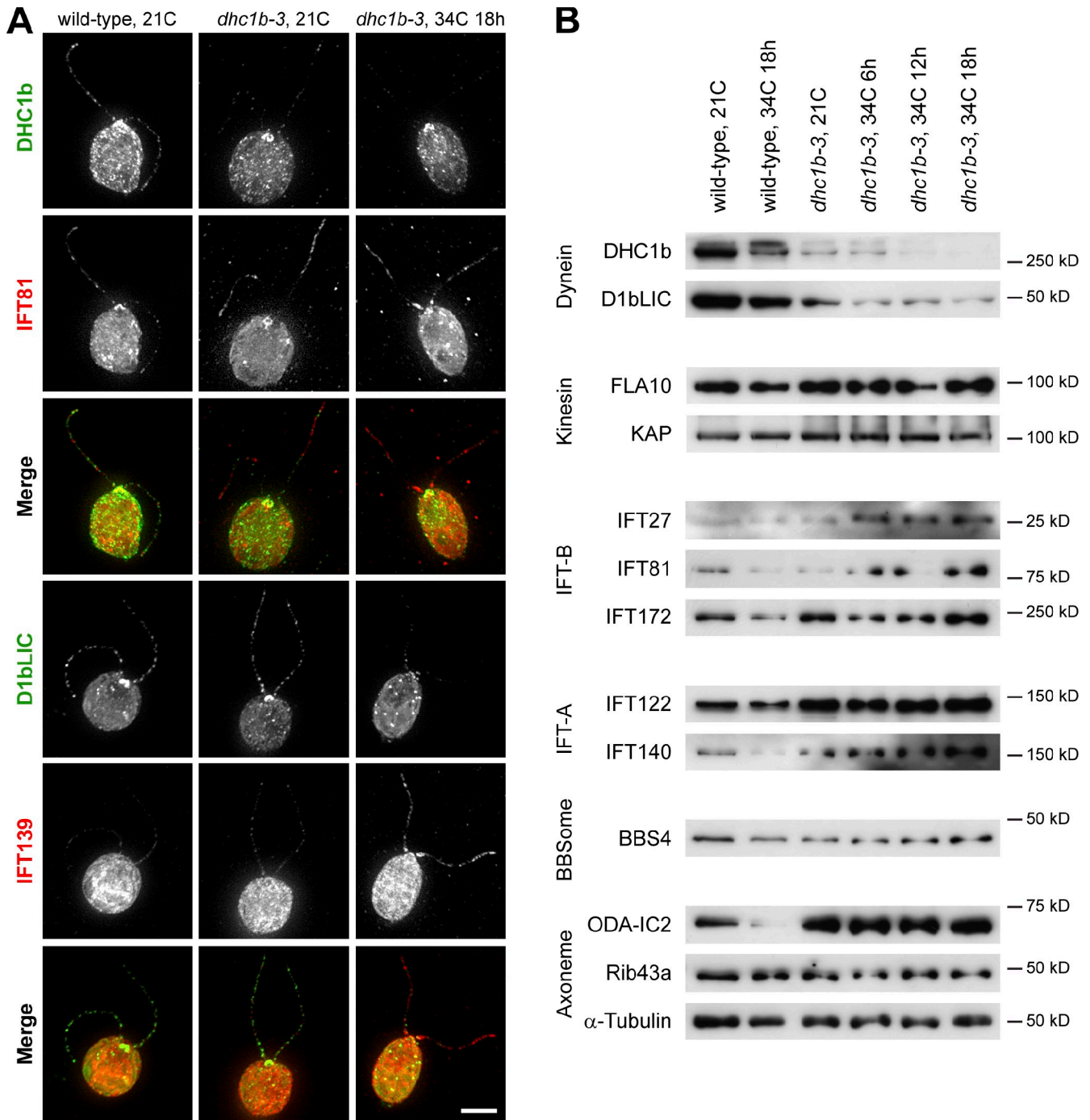


Figure 5. **Accumulation and depletion of flagellar proteins in wild-type and *dhc1b-3* cells at 21 and 34°C.** (A) Immunofluorescence images of wild-type cells at 21°C (left column), *dhc1b-3* cells at 21°C (middle column), and *dhc1b-3* cells after 18 h at 34°C (right column), showing the relationship between dynein depletion and IFT protein accumulation. The top merged images are DHC1b (green) and IFT81 (red). The bottom merged images are D1bLIC (green) and IFT139 (red). Bar, 5 μ m. (B) Western blots of flagella isolated from wild-type and *dhc1b-3* cells at 21°C and increasing time at 34°C, as indicated. Dynein depletion correlated with the accumulation of IFT-B and IFT-A proteins, while the levels of kinesin, BBSome, and many axonemal proteins are unaffected (quantified in Table S2).

Curiously, the depletion of dynein did not cause significant flagellar accumulation of kinesin proteins FLA10 and KAP. This result implies that, unlike IFT proteins, kinesin may not rely on dynein for transport out of flagella. Such a hypothesis is consistent with the absence of KAP-GFP retrograde traces in wild-type cells (Fig. 2 A; Engel et al., 2009a,b), immunofluorescence data showing that FLA10 remains concentrated at the peri-basal body

region of *dhc1b-1* cells in contrast to IFT protein redistribution into swollen stumpy flagella (Pazour et al., 1999a), and Western blotting evidence that FLA10 and KAP do not accumulate in flagella in the absence of D1bLIC (Pedersen et al., 2006). Conversely, in *Caenorhabditis elegans* KAP-GFP has been observed to undergo retrograde transport and to accumulate at the ciliary tips of DHC1b mutants (*che-3*; Signor et al., 1999).

It is also interesting to note that despite the greatly reduced levels of DHC1b protein in *dhc1b-3* flagella at 21°C (Fig. 5 B), retrograde IFT speed and frequency were only moderately decreased (Fig. 2, C and D) and the kinetics of flagellar assembly remained comparable to wild type (Fig. S4). This discrepancy indicates that there is normally an excess of dynein in the flagellum, and begs the question of what function this extra dynein protein is performing. Perhaps it is simply surplus to ensure robust IFT, or perhaps it plays a role in specialized processes such as signal transduction, which our assays do not address.

2D-DIGE proteomic analysis of protein accumulation and depletion in *dhc1b-3* flagella

To get a more comprehensive and unbiased view of how defective retrograde IFT affects the abundance of flagellar proteins, we took a comparative proteomic approach using 2D-DIGE (two-dimensional difference in gel electrophoresis; Viswanathan et al., 2006). Isolated flagella from wild-type cells at 21°C, *dhc1b-3* cells at 21°C, and *dhc1b-3* cells after 18 h at 34°C were each labeled with a different fluorescent dye and run together on a 2D acrylamide gel (Figs. 6 A and S5 A). Spots with the greatest changes in abundance between the three samples were picked for identification via mass spectrometry. The majority of the proteins identified by this 2D-DIGE analysis proved to be known components of the flagellar proteome (Table S3; Pazour et al., 2005). Although numerous spots were proteins of known function, many other proteins were uncharacterized, including a wide assortment of FAPs (flagellar associated proteins; Pazour et al., 2005). The identified spots were clustered into classes based on their relative accumulation or depletion between the three samples (Fig. 6, B and C; Table 1). Proteins that group together in the same class may be transported together by IFT or share similar functions. For example, IFT complex B proteins cluster into class D1, IFT complex A proteins group into class A3, and class A2 is rich in axonemal proteins that may be accumulated turnover products. The potential relationships between the proteins in each class are further explored in the Discussion.

Although many spots were identified as duplicate hits for the same proteins (Table S3), some proteins had spots that grouped into more than one class (Fig. 6 D), indicating that these proteins may receive modifications corresponding to different stages of transport or incorporation into the axoneme. BUG10, Rib43a, and IFT81 all have one spot that clusters with IFT-B proteins (class D1), but also have additional spots with acidic shifts to their isoelectric points (pI) that could indicate phosphorylation. It is plausible that such a modification could be added to these proteins after they have completed anterograde transport. As CAH6 has different spots that group with both IFT-B (class D1) and IFT-A (class A3) proteins, we may have identified forms of CAH6 that are trafficked by each phase of IFT. Interestingly, the transition from IFT-B to IFT-A grouping appears to be accompanied by a slight decrease in CAH6 molecular weight (MW). We identified four FAP103 spots that become increasingly acidic as the spots go from depletion in *dhc1b-3* flagella primarily at 21°C, to depletion at both 21 and

34°C, and finally depletion at only 34°C. These changes in pI are likely related to the protein's function as a nucleoside-diphosphate kinase. Although it is interesting that the activity of FAP103 could be linked to IFT, without further characterization of the protein's role in flagella we cannot speculate about what this connection could be.

Phototaxis, swimming speed, and flagellar beat frequency are altered in *dhc1b-3* cells

Although our results thus far indicate that fully functional retrograde IFT is not required for the maintenance of flagellar length, the changes in *dhc1b-3* flagellar protein composition suggest that retrograde IFT may have an influence on flagellar function. We therefore tested several flagellar functions to see if they are affected by the reduction of retrograde IFT. In dish-based (Fig. 7 A) and single-cell (Fig. 7 B) phototaxis assays at both 21 and 34°C, wild-type cells swam toward the light source and *agg1* cells (Smyth and Ebersold, 1985) swam away from the light. However, *dhc1b-3* cells showed a switch from negative to positive phototaxis when shifted from 21 to 34°C (Fig. 7, A and B). This switch in phototactic sign was seen after 6 h at 34°C in the dish assays and after 12 h at 34°C in the single-cell measurements, a disparity that may be due to intrinsic differences between the assays. Bulk cell migration in the dish assay was the result of 15 min of continuous light exposure, whereas single-cell measurements were acquired 30 s after turning on the light source, before cells could accumulate at one side on the dish. Collectively, it may be inferred that *dhc1b-3* cells incubated at 34°C for 6 h are at an intermediate stage in the switch from negative to positive swimming direction, and that by 12 h at 34°C they exhibit strong positive phototaxis. Because *dhc1b-3* cells at 21°C showed the opposite phototaxis response to the wild-type strain from which they were generated, moderate reductions in retrograde IFT that do not affect anterograde IFT or flagellar length may still significantly influence cell signaling. Mutations in the BBSome component BBS4 have also been shown to affect phototaxis (*bbs4-1*; Lechtreck et al., 2009). Although reduced retrograde IFT in *dhc1b-3* cells would also impair traffic of the BBSome, the *dhc1b-3* and *bbs4-1* phototaxis defects are distinct. Instead of switching the sign of phototaxis, *bbs4-1* cells swam in random directions, irrespective of temperature. However, this difference may be due to the fact that the *bbs4-1* mutant was generated from an *agg1* strain.

The swimming speed of *dhc1b-3* cells was normal at 21°C, but was greatly diminished after 12 h at 34°C (Fig. 7 C). Furthermore, the beat frequency of *dhc1b-3* flagella was decreased at both 21 and 34°C (Fig. 7 D), a telltale sign of an axonemal outer dynein arm (ODA) defect. Because this drop in beat frequency manifests in permeabilized ATP-reactivated cell models in addition to live cells, it is likely caused by a structural defect in the axoneme (although we did not identify an obvious deficiency in TEM cross sections of mutant axonemes; unpublished data). Thus, it appears that retrograde IFT helps maintain both physical (beat frequency) and signaling (phototaxis sign) functions of *Chlamydomonas* flagella. This is likely related to the turnover of flagellar proteins, a connection that we see some evidence for in the 2D-DIGE results

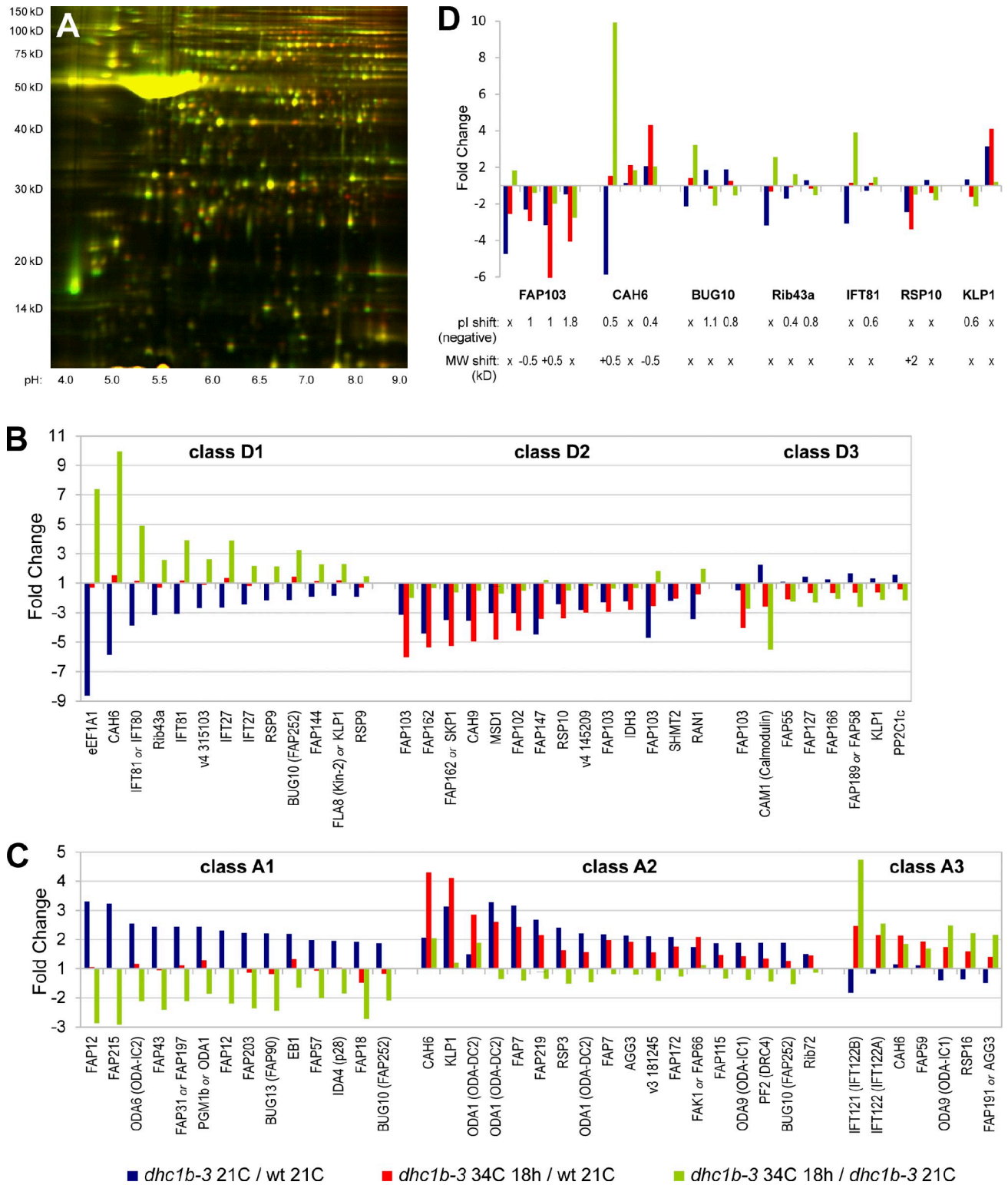


Figure 6. **2D-DIGE proteomic analysis of flagellar protein accumulation and depletion in *dhc1b-3* cells at 21 and 34°C.** (A) Fluorescence image of the 2D-DIGE analytical gel (*dhc1b-3* 21C = green; *dhc1b-3* 34C 18 h = red; for overlays of all three samples see Fig. S5). The large yellow spot at ~50 kD is tubulin. (B and C) Depletion and accumulation profiles for every spot that was identified by mass spectrometry. Fold change is a spot's relative fluorescence between two samples on the analytical gel (1 = equal abundance for both samples). Class D1 = depleted in only *dhc1b-3* 21C vs. wild-type 21C (accumulated in *dhc1b-3* 34C 18 h vs. *dhc1b-3* 21C); class D2 = depleted in both *dhc1b-3* 21C vs. wild-type 21C and *dhc1b-3* 34C 18 h vs. wild-type 21C; class D3 = depleted in only *dhc1b-3* 34C 18 h vs. wild-type 21C (depleted in *dhc1b-3* 34C 18 h vs. *dhc1b-3* 21C); class A1 = accumulated in only *dhc1b-3* 21C vs. wild-type 21C (depleted in *dhc1b-3* 34C 18 h vs. *dhc1b-3* 21C); class A2 = accumulated in both *dhc1b-3* 21C vs. wild-type 21C and *dhc1b-3* 34C 18 h vs. wild-type 21C; class A3 = accumulated in only *dhc1b-3* 34C 18 h vs. wild-type 21C (accumulated in *dhc1b-3* 34C 18 h vs. *dhc1b-3* 21C). (D) The profiles of proteins identified in multiple spots that group into more than one class, suggesting IFT-related protein modifications. Negative pl shifts (minus sign omitted) and MW shifts, both relative to the spot with the highest pl, are noted.

Table 1. Flagellar proteins with changing abundance identified by 2D-DIGE analysis

	21°C only	21 and 34°C	34°C only
<i>dhc1b-3</i> depletion vs. wt	Class D1 (mut-34A): eEF1A1, CAH6, IFT80, Rib43a, IFT81, v4 315103, IFT27, RSP9, BUG10 (FAP252), FAP144, FLA8 or KLP1	Class D2 (mut-34NC): FAP103, FAP162, FAP102, FAP147, RSP10, v4 145209, RAN1	Class D3 (mut-34D): FAP103, CAM1 (calmodulin), FAP55, FAP127, FAP166, FAP189 or FAP58, KLP1, PP2C1c
<i>dhc1b-3</i> accumulation vs. wt	Class A1 (mut-34D): FAP12, FAP215, ODA6 (ODA-IC2), FAP43, FAP31 or FAP197, PGM1b or ODA1, FAP203, BUG13 (FAP90), EB1, FAP57, IDA4 (p28), FAP18, BUG10 (FAP252)	Class A2 (mut-34NC): CAH6, KLP1, ODA1 (ODA-DC2), FAP7, FAP219, RSP3, AGG3, FAP172, FAK1 or FAP66, FAP115, ODA9 (ODA-IC1), PF2 (DRC4), BUG10 (FAP252), Rib72	Class A3 (mut-34A): IFT121, IFT122, CAH6, FAP59, ODA9 (ODA-IC1), RSP16, FAP191 or AGG3
No change	RSP10, Rib43a, FAP164, ICL1, ODA13 (ODA-LC6), IDA5 (actin), IFT81, FAP161, IFT88, FAP52		

Proteins are grouped by relative accumulation or depletion in *dhc1b-3* flagella at 21 and 34°C vs. wild-type flagella at 21°C (as plotted by blue and red bars in Fig. 6). Duplicate hits, tubulin, and proteins not found in the flagellar proteome have been excluded from this list. For the complete catalogue and description of analyzed 2D-DIGE spots, see Fig. S5 and Table S3. Classes D1 and A3 are proteins accumulated in *dhc1b-3* flagella after 18 h at 34°C vs. *dhc1b-3* flagella at 21°C (as plotted by green bars in Fig. 6, indicated on the table by “mut-34A”). Classes D3 and A1 are proteins depleted in *dhc1b-3* flagella after 18 h at 34°C vs. *dhc1b-3* flagella at 21°C (indicated by “mut-34D”). Proteins in classes D2 and A2 experience no change in flagellar abundance when *dhc1b-3* cells are shifted from 21 to 34°C (indicated by “mut-34NC”). Proteins are listed in descending order of fold accumulation or depletion (except the “No change” class).

(see Discussion). It will be interesting to determine whether additional flagellar functions, such as mating, are impacted by the *dhc1b-3* mutation.

Discussion

2D-DIGE proteomics reveals distinct classes of IFT-related flagellar proteins

Class D1 proteins accumulate in *dhc1b-3* flagella at 34°C vs. 21°C (Fig. 6, green bars), and include the IFT complex B proteins IFT81, IFT80, and IFT27. The increase in IFT81 and IFT27 abundance is consistent with our immunofluorescence and Western blot measurements (Fig. 5). Other proteins that group into the D1 profile may interact with IFT complex B as obligate cargos, including uncharacterized proteins FAP144, BUG10, and v4 315103. Of particular interest is eukaryotic translation elongation factor 1A1 (eEF1A1), a small GTPase that is involved in a diverse array of cellular processes in addition to its canonical function of delivering aa-tRNA to the ribosome (Mateyak and Kinzy, 2010). The 100-kD spot was double the expected MW for eEF1A1 and was also a 100% confidence hit for α -tubulin. Whereas tubulin hits for other spots in this study are less compelling because they are near tubulin's expected 50 kD, this spot was well away from the tubulin region of the gel, at a MW and pI predicted for an eEF1A1-tubulin dimer. Although it seems unlikely that such a dimer would survive denaturing electrophoresis, eEF1A does bind directly to microtubules (Moore and Cyr, 2000). eEF1A has been shown in different studies to bundle, stabilize, and sever microtubules (Durso and Cyr, 1994; Shiina et al., 1994; Moore et al., 1998). Although these diverse observations do not paint a clear picture of eEF1A1 in flagella, it is enticing for flagellar length control models that a small GTPase regulator of both translation and microtubule dynamics may be an obligate cargo of anterograde IFT.

The other group of proteins that accumulate in *dhc1b-3* flagella at 34°C vs. 21°C is class A3, which includes the IFT complex A components IFT121 (IFT122B, IFTA-1) and IFT122 (IFT122A, DAF-10). Both proteins are required for retrograde transport and IFT122 has been implicated in the ciliary traffic of

signaling proteins, including Gli2, Gli3, and TULP3 (Qin et al., 2001; Blacque et al., 2006; Cortellino et al., 2009; Mukhopadhyay et al., 2010; Ocbina et al., 2011; Qin et al., 2011). Other accumulated proteins in the A2 and A3 classes are likely cargoes that are ordinarily trafficked by retrograde IFT out of the flagella. These proteins include axonemal components such as ODA-DC2 (ODA1), ODA-IC1 (ODA9), KLP1, FAK1, RSP16, RSP3, DRC4 (PF2), FAP7, FAP219, as well as the “membrane plus matrix” proteins CAH6 and AGG3. The accumulation of ODA proteins and FAK1 could be indicative of outer dynein arm defects that may underlie the reduction in flagellar beat frequency (Fig. 7 D). CAH6 appears to be an abundant IFT cargo, with prominent spots in classes associated with IFT-B (D1), IFT-A (A3), and accumulated cargo (A2). As *Chlamydomonas* flagella are the only part of the cell membrane in direct contact with the extracellular environment, CAH6 may be using IFT to facilitate CO₂ metabolism. The loss of ~1 kD as CAH6 transitions from IFT-B– to IFT-A–associated forms could be related to this function. The accumulation of AGG3 is interesting as it may explain the change in *dhc1b-3* phototaxis direction (Fig. 7, A and B). AGG3 RNAi induces the same phenotype as the *agg1* mutant, where cells swim away from light (Iomini et al., 2006). Thus, it is logical that the accumulation of AGG3 in *dhc1b-3* flagella could cause a switch to positive phototaxis.

Class A1 proteins are depleted from *dhc1b-3* flagella at 34°C vs. 21°C, and have the opposite abundance as the IFT-B proteins in class D1. Most of the A1 proteins are uncharacterized (FAP12, FAP215, FAP31, FAP43, FAP203, BUG13, FAP57, BUG10, and FAP18) and many are in the membrane plus matrix fraction. One of the few well-characterized proteins in the group is EB1, which is found at the flagellar tip and stabilizes microtubules (Pedersen et al., 2003; Schröder et al., 2007). As very few factors have been identified that regulate incorporation and removal of axonemal components, it would be of great interest if any of these uncharacterized proteins share EB1's localization at the flagellar tip.

The second group of proteins depleted from *dhc1b-3* flagella at 34°C vs. 21°C is class D3, which also primarily

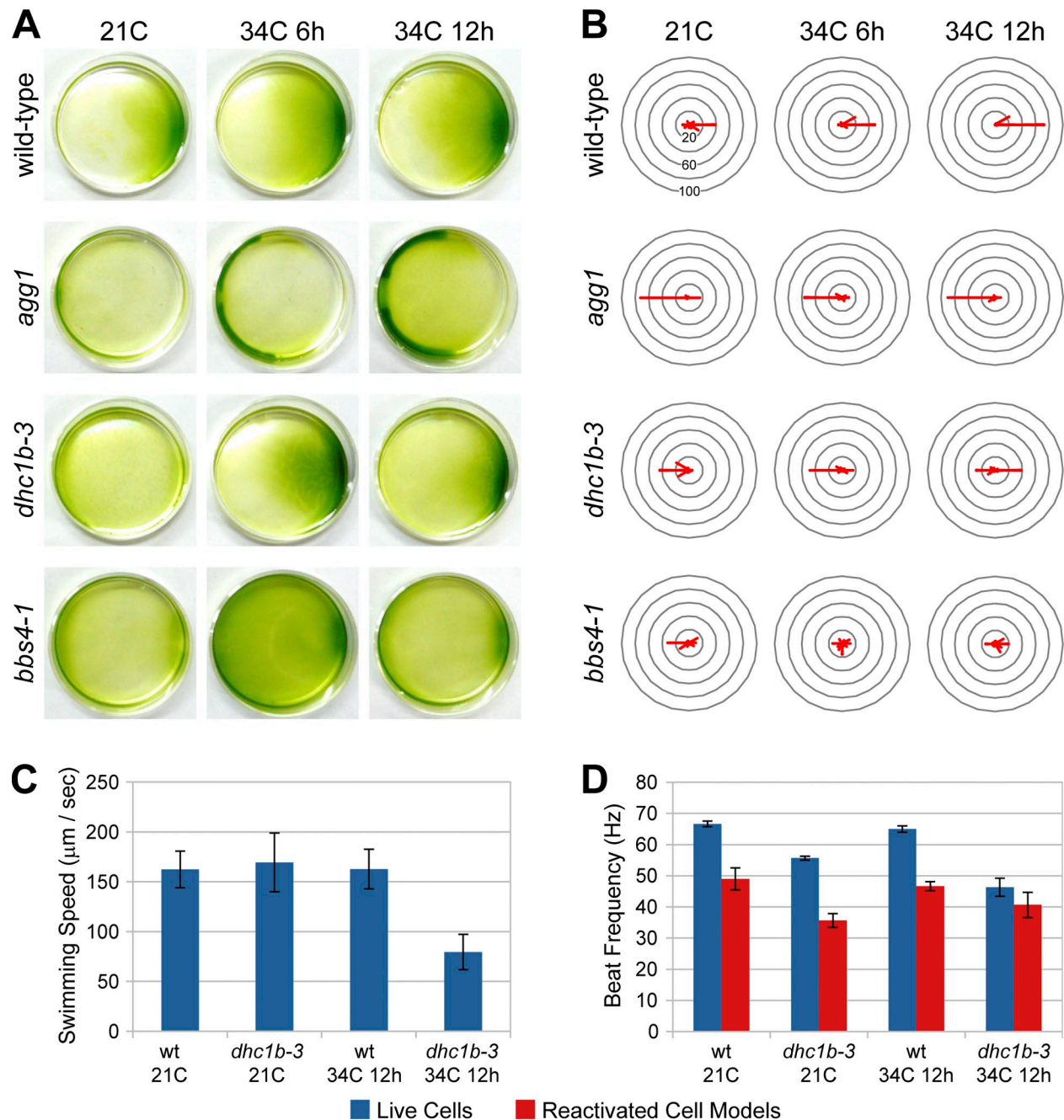


Figure 7. **Effects of *dhc1b-3* on phototaxis, swimming speed, and flagellar beat frequency.** Phototaxis direction of wild-type, *agg1*, *dhc1b-3*, and *bbs4-1* strains at 21 and 34°C, measured by the bulk migration of cells in dish assays (A) and the swimming trajectory of individual cells (B). For A and B, green light illumination is directed from the right. The compass plots in B indicate the percentage of cells swimming in a particular direction relative to the light source, grouped into 30-degree bins. Swimming speed (C) and flagellar beat frequency (D) of wild-type and *dhc1b-3* cells at 21 and 34°C. Reactivated cell models are fixed permeabilized cells that have been treated with ATP, reinitiating flagellar beating (Kamiya and Witman, 1984). Error bars indicate standard deviation.

consists of uncharacterized proteins. Calmodulin has the opposite accumulation profile from IFT121, potentially providing a hint about how calmodulin is transported in flagella. FAP55, FAP127, FAP189/FAP58, and FAP166 are all axonemal proteins with very similar D3 profiles, so perhaps some of these structural proteins interact or even preassemble into a larger complex before being carried into flagella by IFT. FAP103 and FAP162 are repeatedly found among the top hits in classes D2 and D3. These two proteins are exclusively

found in the membrane plus matrix fraction and may rely on DHC1b activity for entry into flagella.

Interestingly, IFT88 clusters into the “no change” class (Table 1). Although this could mean that there is something fundamentally different about the transport of IFT88, it should be noted that IFT81 also has one spot that sees no change in abundance, so we may have simply not picked the IFT88 spot that fits the D1 profile. Regardless, this observation warrants additional investigation.

Table 2. Summary of the *dhc1b-3* IFT and flagellar phenotypes compared to wild-type cells at the same condition

<i>dhc1b-3</i> vs. wild type	Retrograde		Anterograde		Flagellar assembly	Length maintenance	Flagellar shortening
	Speed	Freq.	Speed	Freq.			
	%	%	%	%			
21°C	75	76	102	99	++	++	++
34°C 6 h	47	35	91	63	–	+	ND
34°C 12 h	40	20	89	58	--	+	++

"++" = same phenotype as wt; "+" = close to wt; "–" = strong defect relative to wt; "--" = complete defect; ND = no data.

The relationship between anterograde and retrograde IFT, and implications for flagellar length control

One clear conclusion that can be drawn from our data is that anterograde and retrograde IFT are coupled, but it is not a 1:1 relationship (Table 2). It may be easiest to understand the *dhc1b-3* mutant's flagellar phenotype in respect to this loose coupling. In *dhc1b-3* cells at 21°C, anterograde IFT appears unaffected by a moderate ~25% decrease in retrograde IFT speed and frequency relative to wild-type levels. It is perhaps no surprise then that *dhc1b-3* flagellar assembly at 21°C is indistinguishable from wild type. After incubating *dhc1b-3* cells for 12 h at 34°C, retrograde speed drops to ~40% of wild-type levels and retrograde frequency drops to ~20% of wild type. This strong retrograde IFT defect is coupled to less dramatic changes in anterograde speed and frequency, which decrease to ~90% and ~60% of wild type, respectively. Presumably, this level of anterograde IFT is sufficient to maintain flagella with little shortening, but is insufficient to drive flagellar assembly.

How well does the *dhc1b-3* phenotype equate with the balance-point model of flagellar length control, which asserts that flagellar length is the resultant of opposing assembly and disassembly rates at the flagellar tip (Marshall and Rosenbaum, 2001; Marshall et al., 2005; Avasthi and Marshall, 2012)? The

current iteration of the balance-point model predicts that, with a 40% reduction in anterograde IFT frequency at 34°C, both existing and newly assembling *dhc1b-3* flagella should approach the same intermediate steady-state length (Fig. 8). Experimentally, however, *dhc1b-3* flagellar length at 34°C is bi-stable. Existing flagella only slightly shorten, whereas new flagella experience strong assembly defects. Thus, there are likely uncharacterized intrinsic differences between established full-length flagella and short assembling flagella that are not yet factored into the balance-point model. One possibility is that stabilizing factors at the tips of full-length flagella, such as EB1 (Schröder et al., 2007) or the distal filaments (Dentler and Rosenbaum, 1977; Dentler, 1980), could make the disassembly rate length dependent.

An important caveat about the nearly full-length *dhc1b-3* flagella at 34°C is that we did not quantify IFT train size. If the drop in retrograde IFT leads to a reduction in anterograde train size, then the effect on cargo delivery to the flagellar tip could potentially be very small. Future refinement of the balance-point model will require precise measurements of IFT train size, speed, and frequency, or perhaps even direct quantification of axonemal cargo delivery to the flagellar tip, coupled with single-cell measurements of flagellar length changes.

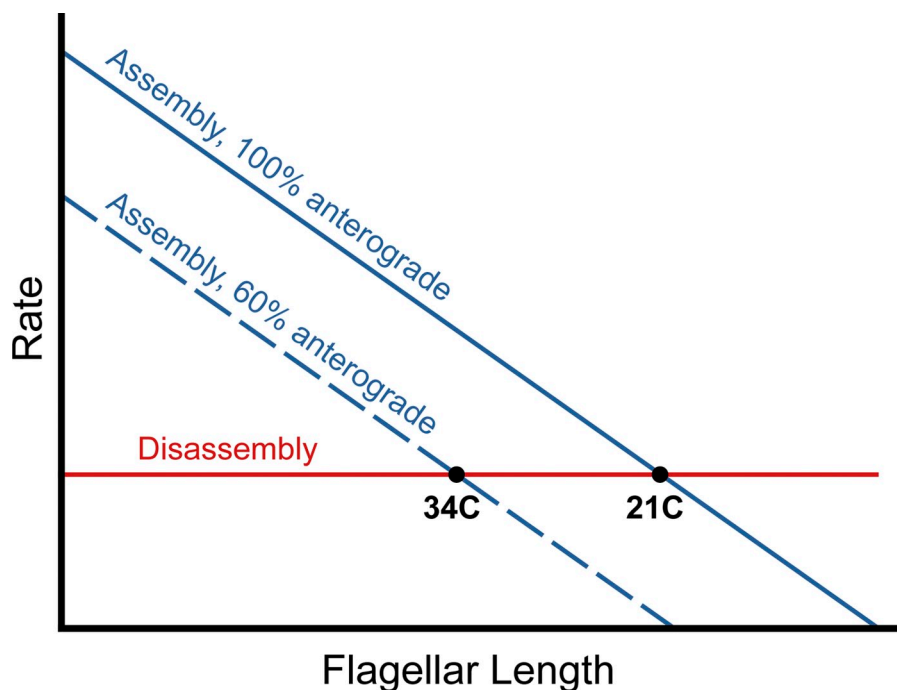


Figure 8. Comparison of the *dhc1b-3* phenotype with the current balance-point model of flagellar length control. The horizontal red line represents the length-independent disassembly rate and the blue lines are length-dependent assembly rates at 21°C (solid line) and 34°C (hashed line, 40% reduction in anterograde IFT frequency). The balance-point model predicts that both existing and assembling *dhc1b-3* flagella should approach the same steady-state length (points indicated by "21C" and "34C"), where the assembly and disassembly rates are equal.

Finally, although our results clearly demonstrate the requirement for DHC1b in flagellar assembly, it remains to be shown why this is the case. The plausible default hypothesis is that continuous anterograde transport requires the recycling of IFT particles by retrograde IFT. However, it is possible that a signaling function of retrograde IFT is the true requirement for anterograde transport initiation. The unknown mechanism underlying the loose coupling between anterograde and retrograde IFT observed in the *dhc1b-3* mutant may provide the key to understanding the reliance of flagellar assembly on retrograde transport.

Conclusion

The *dhc1b-3* mutant's unique kinetics of flagellar loss indicate that robust retrograde IFT is dispensable for the maintenance of flagellar length. However, it is clear that retrograde transport still plays key roles during flagellar assembly and in maintaining flagellar composition and function. Comparative proteomics of *dhc1b-3* flagella revealed several distinct classes of known and uncharacterized IFT-related proteins that warrant further investigation. Like the *fla10* mutant has done for kinesin-driven anterograde transport, the inducible and reversible inhibition of dynein function afforded by the *dhc1b-3* mutant should prove exceptionally useful for future studies of retrograde IFT.

Materials and methods

Strains and culture conditions

Vegetative cells were grown in liquid Tris-acetate-phosphate (TAP) or minimal (M) media (Harris et al., 2009) at 21 or 34°C with constant aeration. Gametogenesis was induced by suspending cells in M-N media (M media without nitrogen). Wild-type mt+ (CC-125), wild-type *agg1* mt- (CC-124), *dhc1b-1* mt+ (CC-3711), and S1C5 mt- (CC-1952) strains were obtained from the Chlamydomonas Resource Center (University of Minnesota, Minneapolis, MN). The KAP-GFP *fla3-1* (CC-4296) strain (Mueller et al., 2005) was provided by M. Porter (University of Minnesota, Minneapolis, MN). The *bbs4-1* mt- (CC-4371) strain (Lehtreck et al., 2009) and the *dhc1b-2* strain (Pazour et al., 1999b; Witman, 2011) were supplied by G. Witman (University of Massachusetts Medical School, Worcester, MA).

Isolation of the *dhc1b-3* mt+ (CC-4422) strain is detailed in Fig. S2 E. Liquid cultures of wild-type CC-125 cells were mutagenized by exposure to UV radiation at a dose sufficient to kill 90% of the cells (LD90). Cells were then embedded in soft agar (0.35% agar in TAP) and grown for 1 wk at 34°C. Punctate colonies were picked out of the agar, grown in TAP at both 21 and 34°C, and checked by live-cell microscopy for *ts* conditional motility phenotypes. After isolation, the *dhc1b-3* mt+ strain was backcrossed twice; the mutant phenotype consistently segregated 1:1. The *dhc1b-3* mt- (CC-4423) strain was generated in the course of tetrad analysis for linkage mapping. The *dhc1b-3 fla10* (CC-4424) double mutant was produced by mating *dhc1b-3* mt+ (CC-4422) to *fla10-1* mt- (CC-1919).

To produce diploid strains, electroporation was used to transform *dhc1b-1* mt+ (CC-3711) cells with the pHyg3 hygromycin (hyg) resistance vector and *dhc1b-3* mt- (CC-4423) cells with the pS1103 aphVIII paromycin (par) resistance vector (plasmids were obtained from the Chlamydomonas Resource Center). The two drug-resistant strains were then mated, plated on 1.5% agar hyg/par TAP plates, and grown at room temperature in constant light until colonies appeared. Colonies were picked, struck onto 1.5% agar hyg/par TAP plates, and tested by PCR for mating type and the presence of the *dhc1b-3* point mutation (Fig. S2 D). Positives were restripped onto new plates and retested by PCR.

Linkage mapping and sequencing the *dhc1b-3* mutation

To locate the genomic region harboring the *dhc1b-3* mutation, the mutant (mt+, CC-4422) was crossed to the S1C5 polymorphic strain (mt-, CC-1952; Gross et al., 1988) and tetrad progeny were isolated by dissection (Harris et al., 2009). The recombination frequencies between

dhc1b-3 and the genetic markers SSR157-44 (5'-ACTGGGTGTGCACATACATGGTGG-3', 5'-ACCTCAACCTTGGGGACGTAAAAT-3'), SSR75-62 (5'-CCCCGCTTACCTACCCACTATGCTT-3', 5'-CGACAGTGGATGTGCCAGTAGGA-3'), FA1 (5'-ACGAGGAGGACATTCGGGAGCTGC-3', 5'-GCTCAGCCGTCCAAGGAAGCAATG-3', 5'-GTCCAAGCAGGTCCAAGCCGTCAA-3'; Kathir et al., 2003), and the mating-type locus (MT; 5'-CGACGACTTGGCATCGACAGGTGG-3', 5'-CTCGGCCAGAACCCTTCATAGGTGG-3', 5'-GCTGGCATTCTGTATCCTTGACGC-3', 5'-GCGGCGTAA-CATAAAGGAGGGTCCG-3'; Zamora et al., 2004) were determined by detecting polymorphic PCR products in tetrad progeny with the *dhc1b-3* flagellar assembly defect (Fig. S2 A). Recombinants were found for all markers other than SSR75-62, narrowing down the region where the mutation could be located.

To determine the precise location of the *dhc1b-3* mutation, we sequenced cDNA of the DHC1b gene. Total RNA was isolated from *dhc1b-3* and wild-type cells by TRIzol/Chloroform extraction, then cDNA was synthesized using Superscript III reverse transcription and random hexamer primers (Life Technologies). Using the primer pairs shown in Table S1, 29 overlapping PCR products were amplified from the *dhc1b-3* cDNA, spanning the length of the coding sequence (CDS) and flanking untranslated regions (UTRs). PCR products were processed through a cleanup kit (QIAGEN) and then sequenced (Elim Biopharm). Only a single base pair differed from the wild-type sequence (12313 of the CDS). The mutation was confirmed by sequencing wild-type cDNA spanning the region, and through differential cutting of PCR products with restriction enzymes (Fig. S2, B and C). Noncomplementation in the *dhc1b-1 dhc1b-3* diploid strain proved that the *dhc1b-3* phenotype was due to a mutation in DHC1b (Fig. S2 D).

To examine the evolutionary conservation of the DHC1b C-sequence, the DHC1b amino acid sequences for the nine species displayed in Fig. S3 were aligned using ClustalX (Larkin et al., 2007). Aligned residues were colored to indicate conservation if they met the default thresholds that Clustal uses to assign Zappo colors, which are more stringent for some amino acid types than others. Instead of Zappo colors, the ClustalX "colprot.xml" file was modified to display a red-to-blue gradient corresponding to Kyte-Doolittle hydrophobicity values (Kyte and Doolittle, 1982). Dark blue indicates conserved hydrophilic residues, and dark red amino acids are conserved hydrophobic.

Flagellar regeneration and shortening

Flagellar regeneration was initiated by pH shock, as described previously (Lefebvre, 1995). TAP cell cultures were decreased to pH 4.5 (with 0.5 M acetic acid), incubated for 1.5 min to trigger flagellar abscission, then returned neutral pH (with 0.5 M KOH) to enable regeneration of flagella. To examine flagellar shortening, cells were grown in M media, incubated for 12 h at 21 or 34°C, and then treated with 20 mM NaPPi. Cells were incubated at 21 or 34°C after NaPPi treatment, fixed with glutaraldehyde, and imaged with a DeltaVision microscope (Applied Precision). Flagellar lengths were measured using ImageJ software (National Institutes of Health, Bethesda, MD).

Live-cell measurements of IFT by DIC and TIRF microscopy

Live-cell microscopy of IFT and kymograph analysis were performed as detailed previously (Engel et al., 2009a,b). For TIRF, 15 μ l of GFP-expressing cells in TAP media were mounted on untreated cover glass (22 \times 22 mm; Corning), inverted onto a slide (Gold Seal; Thermo Fisher Scientific) with a petroleum jelly spacer. Movies were acquired at \sim 30 frames per second at room temperature (for <1 h after removing cells from 34°C) with NIS-Elements software (Nikon) and a cooled EM CCD camera (QuantEM:512SC; Photometrics) attached to a microscope (Eclipse Ti; Nikon) equipped with a 100 \times /1.49 NA TIRF oil immersion objective, a built-in perfect focus system, and through-the-objective TIRF with motorized control of the incident angle of illumination.

DIC imaging was performed essentially as described previously (Craig et al., 2010). Cells were immobilized between the cover glass and a 0.5% agarose pad. Movies were acquired at \sim 24 frames per second at room temperature (for <1 h after removing cells from 34°C) with NIS-Elements software (Nikon) and a Clara Interline camera (Andor Technology) attached to a microscope (Eclipse Ti; Nikon) equipped with a 100 \times /1.4 NA TIRF oil immersion objective, a 1.4 NA oil immersion condenser, and a manually operated translating rotating stage, which enabled the specimen to be rotated so that flagella were perpendicular to the axis of DIC shear before imaging.

Kymographs for both the TIRF and DIC movies were generated with MetaMorph software (Molecular Devices) by drawing a line along the flagella with a width slightly greater than the width of the flagella.

The maximum intensity value (for TIRF) or average intensity (for DIC) for each point along the length of the line was plotted for each frame in the movie. IFT train speed and frequency were quantified from the kymographs by measuring the slope and number of traces, respectively.

Western blotting and immunofluorescence microscopy

Flagella for Western blots were isolated as described previously (Witman et al., 1972) by pH shock, pelleting the cell bodies (1,000 rcf; 5 min), centrifuging the supernatant onto an underlayered 25% sucrose cushion (2,500 rcf; 10 min, no brake), collecting the interface plus supernatant, and then pelleting the flagella (10,000 rcf; 20 min). Western blots were quantified by calculating the background-normalized integrated intensity using ImageJ. Immunofluorescence was performed essentially as described in Keller et al. (2010). In brief, cells were plated on polylysine-coated cover glass, fixed in methanol at -20°C , blocked in 5% BSA, 1% fish gelatin, and 10% normal goat serum in PBS, and incubated with primary antibodies overnight (1:100). After several washes with PBS, the cells were incubated with secondary antibodies (1:200; Jackson ImmunoResearch Laboratories, Inc.), washed with more PBS, incubated with 1 $\mu\text{g}/\text{ml}$ DAPI, and mounted on slides with Vectashield mounting medium (Vector Laboratories). Cells were imaged using a DeltaVision deconvolution system (Applied Precision) with a 100 \times /1.4 NA oil immersion objective. Z-stacks were acquired, deconvolved, and then made into maximum intensity projections.

The primary antibodies used in this study were generously provided by D. Cole (rat α -KAP, mouse α -FLA10N, mouse α -IFT81, mouse α -IFT172, mouse α -IFT139, rabbit α -IFT140, and rabbit α -IFT122; Cole et al., 1998; Mueller et al., 2005; Behal et al., 2012), George Witman (rabbit α -DHC1b, rabbit α -D1bLIC, and rabbit α -BBS4; Pazour et al., 1999a; Hou et al., 2004; Lechtreck et al., 2009), Joel Rosenbaum (rabbit α -IFT27; Qin et al., 2007), and Richard Linck (rabbit α -Rib43a; Norrander et al., 2000). Antibodies to ODA-IC2 (IC69, clone D6168) and α -tubulin (DM1A, clone T6199) were obtained from Sigma-Aldrich.

2D-DIGE analysis

For each condition (wild-type 21C, *dhc1b-3* 21C, *dhc1b-3* 34C 18 h), two liters of cells were grown in TAP with constant light and aeration. Flagella were isolated as described previously (Witman et al., 1972), yielding 100–200 μg of flagellar protein for each sample. Protein concentrations were adjusted using Western blots with fluorescent secondary antibodies (measured by an Odyssey imager [LI-COR Biosciences]) to ensure equal α -tubulin loading between the samples. The samples were then sent to Applied Biomics, Inc. (Hayward, CA) for 2D-DIGE comparative proteomics.

The three samples were each labeled with a different cy-dye and then run together on a single 2D acrylamide gel (~ 50 μg total protein), where they were separated by isoelectric focusing (isoelectric point, pI) followed by electrophoresis (molecular weight, MW). This “analytical” gel was scanned for the fluorescence signal of each cy-dye (Typhoon imager; GE Healthcare), and DeCyder software (GE Healthcare) was used to perform “in-gel” comparative analysis of every spot. Spots were chosen for mass spectrometry identification based on high relative accumulation or depletion between the three samples. A second “preparative” 2D gel was run with higher protein concentration (~ 200 μg total protein) and spots were picked out of the gel using an automated Ettan Spot Picker (GE Healthcare). For spot identification, picked gel samples were washed, dried, and rehydrated in trypsin digestion buffer at 37°C . Digested peptides were extracted from the gel with trifluoroacetic acid buffer, desalted using C-18 Zip-tips (EMD Millipore), mixed with α -cyano-4-hydroxycinnamic acid matrix (CHCA), and spotted into wells of a matrix-assisted laser desorption ionization (MALDI) plate. The mass spectra (MS) of the peptides in each sample was determined by time of flight (MALDI-TOF) using an Applied Biosystems proteomics analyzer (Life Technologies). The most abundant peptides in each sample were subjected to further fragmentation and tandem mass spectrometry (MALDI-TOF/TOF, MS/MS). Protein identities were determined by searching the *Chlamydomonas* genome sequence (Joint Genome Institute, assembly v4.0) with the combined MS and MS/MS spectra using GPS Explorer software equipped with the MASCOT search engine (Matrix Science, Inc.). Only strong hits with close to 100% confidence were accepted.

Cell motility and phototaxis assays

The phototactic behavior of *Chlamydomonas* cells was observed and quantified as described previously (Wakabayashi et al., 2011). For dish assays, cells were washed with buffer (5 mM Hepes, pH 7.4, 0.2 mM

EGTA, 1 mM KCl, and 0.3 mM CaCl_2) and put in Petri dishes. Cell suspensions were illuminated from one side (from the right in Fig. 7) with a green LED ($\lambda = 525$ nm, ~ 50 $\mu\text{mol photons m}^{-2}\text{s}^{-1}$) for 15 min and then photographed. For single-cell analysis, the Petri dishes were placed on the stage of an inverted microscope (model IX70; Olympus) and illuminated with the green LED. 30 s after the LED was turned on, the behavior of the cells was video-recorded using a CCD camera (model XC-75; Sony). The angle (θ) between the light direction and the swimming direction was measured from the tracks of swimming cells using Image Hyper software (Science-Eye).

Reactivation of demembrated cell models was performed by modifying the method described in Kamiya and Witman (1984). After cells were washed with a buffer containing 10 mM Hepes, pH 7.4, and 4% sucrose, cells were demembrated with HMDEKP buffer (30 mM Hepes, 5 mM MgSO_4 , 50 mM potassium acetate, and 1% polyethylene glycol Mr = 20,000) containing 0.1% Igepal CA-630. For reactivation, cell models were suspended in HMDEKP buffer containing 70 U/ml creatine kinase, 5 mM phosphocreatine, and 1 mM ATP.

Flagellar beat frequency was measured as described previously (Kamiya, 2000; Wakabayashi and King, 2006). In brief, a photomultiplier with a gradient filter was attached to a dark-field microscope (model BH-2; Olympus) and cells were observed with dim red light. Cell body vibration signals were transferred to a computer via an A/D converter (model SE-U55X; Onkyo Corporation) and put through a fast-Fourier transform (FFT) using SIGVIEW software (Signalab). Transformed signals were averaged for 30 s. The resulting power spectrum represents the beat frequency distribution, with the peak value indicating the median beat frequency. Swimming velocities were calculated from the tracks left by swimming cells during the extended exposure of images, obtained using Image Hyper software.

Online supplemental material

Fig. S1 shows the phenotype of *dhc1b-2*. Fig. S2 diagrams the scheme used to isolate *ts* motility mutants, and shows the linkage mapping of *dhc1b-3*, the PCR assay used to track the *dhc1b-3* mutation, and noncomplementation in the *dhc1b-1 dhc1b-3* diploid strain. Fig. S3 illustrates the location of the *dhc1b-3* mutation and the conservation of the sequence where it is found. Fig. S4 graphs the kinetics of *dhc1b-3* flagellar loss in synchronized dividing cells and the kinetics of flagellar regeneration in *dhc1b-3* cells at 21°C after increasing preincubation times at 34°C . Together, Fig. S5 and Table S3 detail the complete results from the 2D-DIGE proteomic analysis. Table S1 lists the primers used to sequence the DHC1b cDNA. Table S2 shows the quantification of Fig. 5 B. Videos 1–6 were used to generate the kymographs in Fig. 2, A and B. Online supplemental material is available at <http://www.jcb.org/cgi/content/full/jcb.201206068/DC1>.

We extend a special thanks to Ritsu Kamiya for hosting a two-month stay at the University of Tokyo and to all the members of the Kamiya laboratory for their help and gracious hospitality. We are grateful to Douglas Cole, Dennis Diener, Carlo Iomini, Karl Lechtreck, Richard Linck, and Joel Rosenbaum for generously sharing strains and antibodies. We thank Kurt Thorn and the Nikon Imaging Center at UC San Francisco for assistance with the TIRF microscope.

This work was funded by National Institutes of Health grant R01 GM 097017 (to W.F. Marshall), the Genentech Graduate Fellowship (to B.D. Engel), and the NSF East Asia and Pacific Summer Institutes Program (to B.D. Engel), co-funded by the Japan Society for the Promotion of Science.

Author contributions: B.D. Engel isolated the *dhc1b-3* strain, sequenced the mutation, performed initial characterization of flagellar loss and regeneration, imaged and quantified IFT, and implemented the 2D-DIGE analysis with Applied Biomics, Inc. H. Ishikawa performed the Western blot analysis of flagellar protein levels. K.A. Wemmer created and characterized the *dhc1b-1 dhc1b-3* diploid strain. S. Geimer performed the TEM sample preparation and imaging. K.-i. Wakabayashi measured phototaxis, cell swimming speed, and flagellar beat frequency. M. Hirono performed linkage mapping of the *dhc1b-3* strain and isolated a mating-type minus recombinant used to produce the diploid. B. Craige helped with the DIC microscopy of IFT. G.J. Pazour isolated and characterized the *dhc1b-2* strain. G.B. Witman provided DIC imaging facilities. R. Kamiya aided with TEM and dark-field microscopy, and provided facilities. W.F. Marshall provided facilities and measured flagellar regeneration and shortening. B.D. Engel and W.F. Marshall wrote the paper.

Submitted: 18 June 2012

Accepted: 30 August 2012

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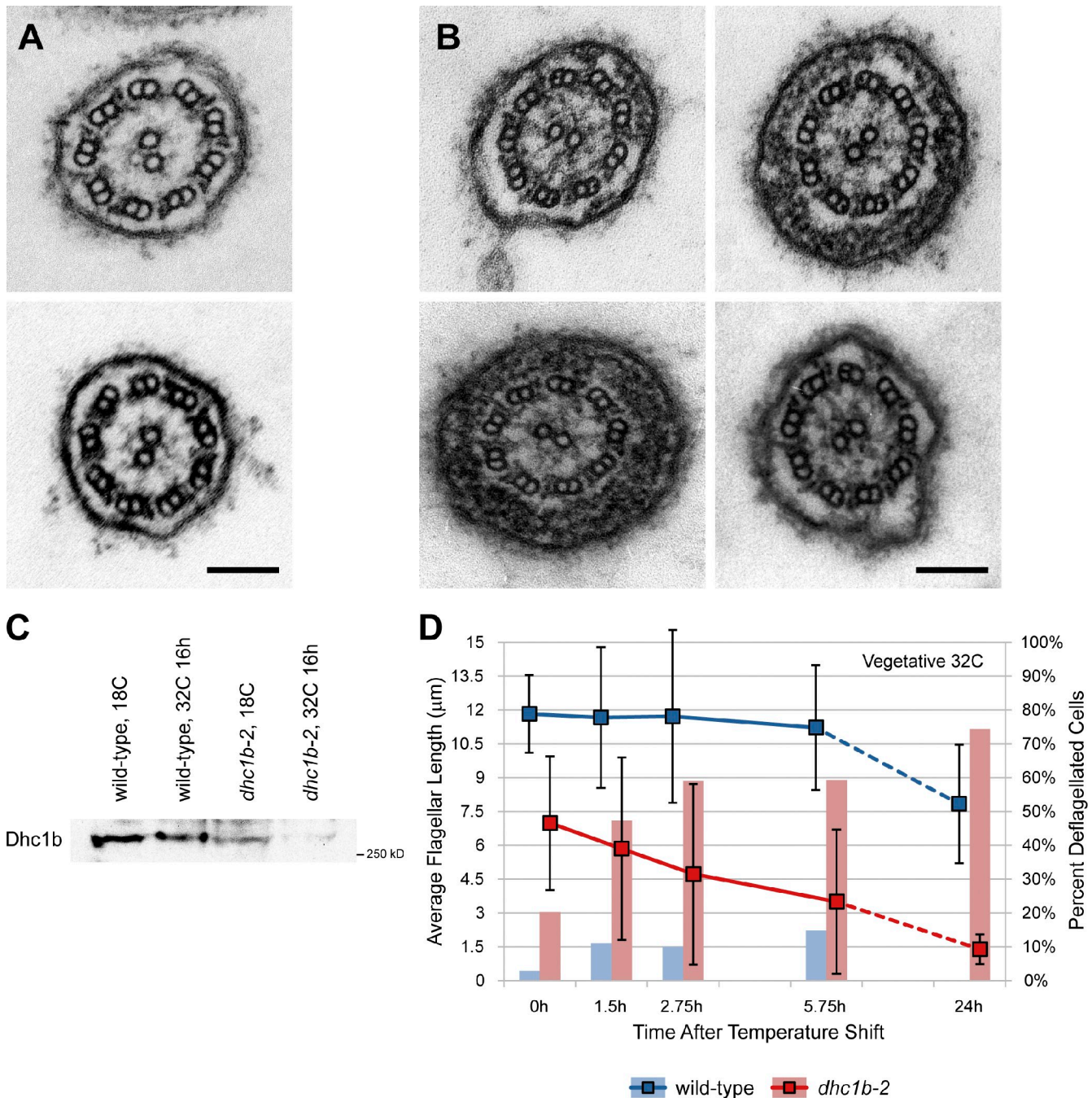
Engel et al., <http://www.jcb.org/cgi/content/full/jcb.201206068/DC1>

Figure S1. **Temperature-sensitive phenotype of the *dhc1b-2* mutant.** Transmission electron microscopy (TEM) cross sections of flagella from wild-type (A) and *dhc1b-2* (B) cells grown at 18°C. Even at this permissive temperature, *dhc1b-2* flagella show a significant accumulation of IFT proteins (electron-dense material between the axoneme and the flagellar membrane). Bars, 100 nm. (C) DHC1b protein levels in whole-cell lysate from wild-type and *dhc1b-2* cells grown at 18°C and after a shift to 32°C for 16 h. DHC1b is reduced in *dhc1b-2* cells at the permissive temperature and further depleted at the restrictive temperature. (D) The kinetics of flagellar shortening and loss in *dhc1b-2* cells at 32°C, compared with a wild-type control. Bar graphs depict the percentage of deflagellated cells, and line graphs show the average flagellar length of flagellated cells. $N_{\text{cells}} = 2,335$ from a single experiment. Unlike *dhc1b-3* (see Fig. 3), *dhc1b-2* flagella are half-length at 18°C and undergo both flagellar shortening and loss when shifted to the nonpermissive temperature, with kinetics that are similar to anterograde IFT mutants such as *fla10*. The lack of *dhc1b-2* flagellar maintenance at 34°C may be indicative of a difference in stability between assembling and full-length flagella. According to this hypothesis, the short *dhc1b-2* flagella at 21°C have not reached a stable length, and thus disassemble when IFT is further impaired at 34°C.

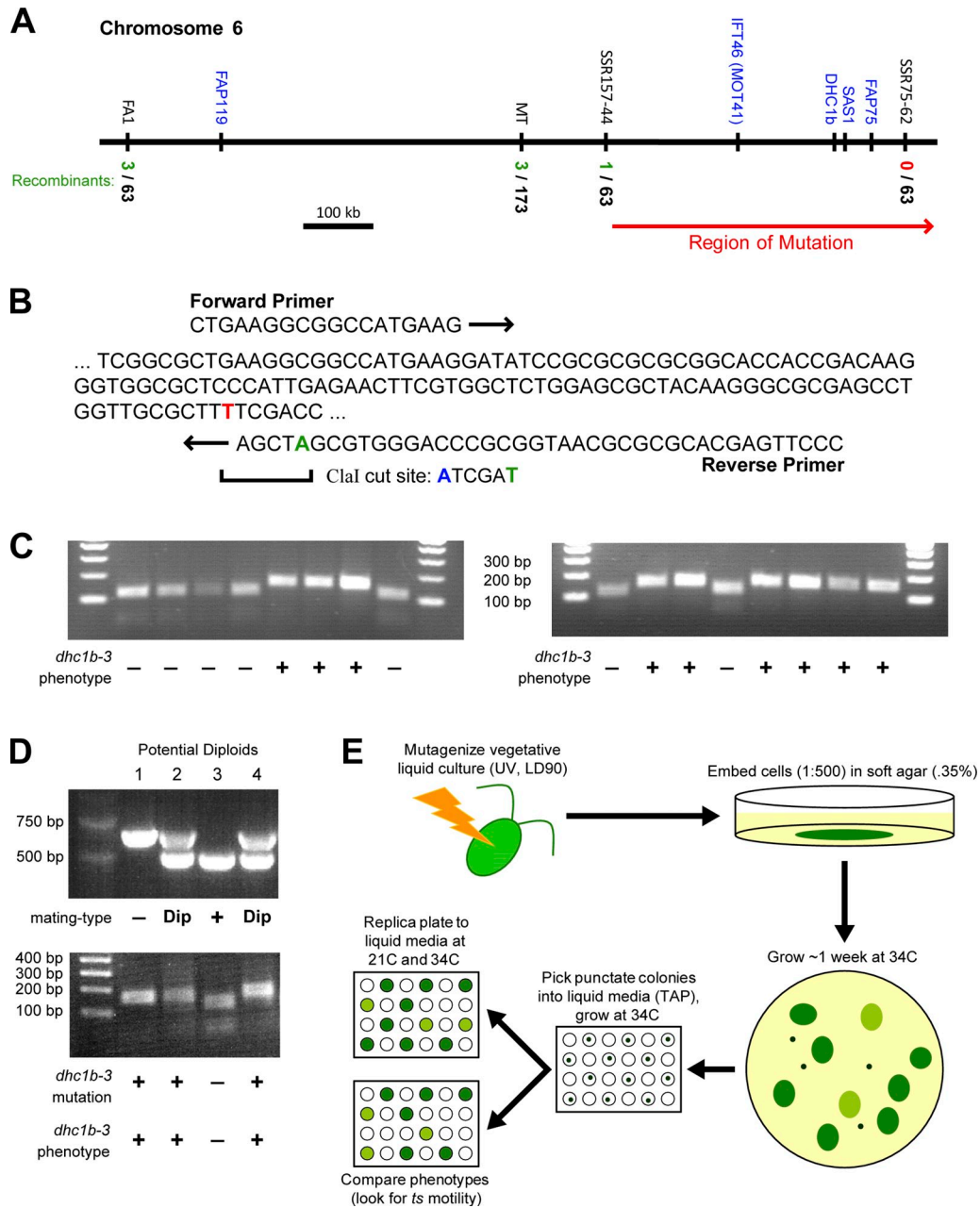


Figure S2. Linkage mapping of the *dhc1b-3* mutant, PCR verification of a single base-pair mutation, *dhc1b-1 dhc1b-3* diploid noncomplementation, and the temperature-sensitive cell motility screening strategy. (A) The mutant phenotype was mapped to the end of chromosome 6 by mating the *dhc1b-3* mutant to polymorphic strain S1C5, performing tetrad dissection, and scoring the recombination frequency of the progeny by PCR-based SNP analysis. Genes listed in blue produce proteins that localize to flagella (Pazour et al., 2005). This analysis also yielded a mating-type minus (*mt-*) *dhc1b-3* strain that was used to produce a diploid strain with the *dhc1b-1* null dynein mutant (Pazour et al., 1999a). (B) PCR scheme for rapid verification of the *dhc1b-3* point mutation. The forward primer is ~100 bp upstream of the mutation, whereas the reverse primer starts immediately after the mutation (in red). The reverse primer has a single mismatch (in green) that creates a *Clal* restriction cut site in PCR products from wild-type (A, in blue) but not mutant (T, in red) genotypes. Because the reverse primer is 40 bp long, cutting the 155-bp PCR product produces a significant band shift when the samples are run on an agarose gel. (C) Using this scheme, raw cell lysates from progeny (in this case, from the *dhc1b-3* × *fla10* mating) were readily screened for the *dhc1b-3* mutation. The plus symbols indicate the presence of the *dhc1b-3* conditional flagellar assembly phenotype, which correlates in each progeny with an undigested PCR product. (D) The *dhc1b-3* mutation does not complement with the *dhc1b-1* null allele in diploid strains. The top gel shows the mating type of four potential diploid strains (Zamora et al., 2004) and the bottom gel shows the presence or absence of the *dhc1b-3* mutation by PCR digestion (same assay as C). The sample in lane 1 is not diploid; its genotype and phenotype are identical to the *dhc1b-3* parental strain (*mt-*, *dhc1b-3* mutation, conditional flagellar loss). The sample in lane 3 is also not diploid; it has the genotype and phenotype of a wild-type *mt+* strain. The samples in lanes 2 and 4 are *dhc1b-1 dhc1b-3* diploid strains with both *mt+* and *mt-* mating-type loci. The diploid strains have *dhc1b-3* mutations and demonstrate conditional flagellar loss at 34°C with similar kinetics to the parental *dhc1b-3* strain. Thus, the *dhc1b-3* and *dhc1b-1* alleles show noncomplementation, confirming that the *dhc1b-3* strain's phenotype is caused by the point mutation in DHC1b. (E) Genetic screen for *ts* motility mutants in *Chlamydomonas*. TAP liquid cultures were exposed to UV light at a dose sufficient to kill 90% of cells (LD90). The cultures were then diluted into low concentration agar (0.35%) TAP plates and grown for 1 wk in fluorescent light at 34°C (the nonpermissive temperature), resulting in embedded clonal colonies. Diluting the UV-treated culture 1:500 in agar resulted in ~1 colony/cm². Although motile colonies became diffuse, colonies with motility defects remained punctate. Punctate colonies were picked out of the soft agar, grown in 96-well plates at 34°C, then replica plated to additional 96-well plates at both 34 and 21°C and examined by microscopy for *ts* motility phenotypes.

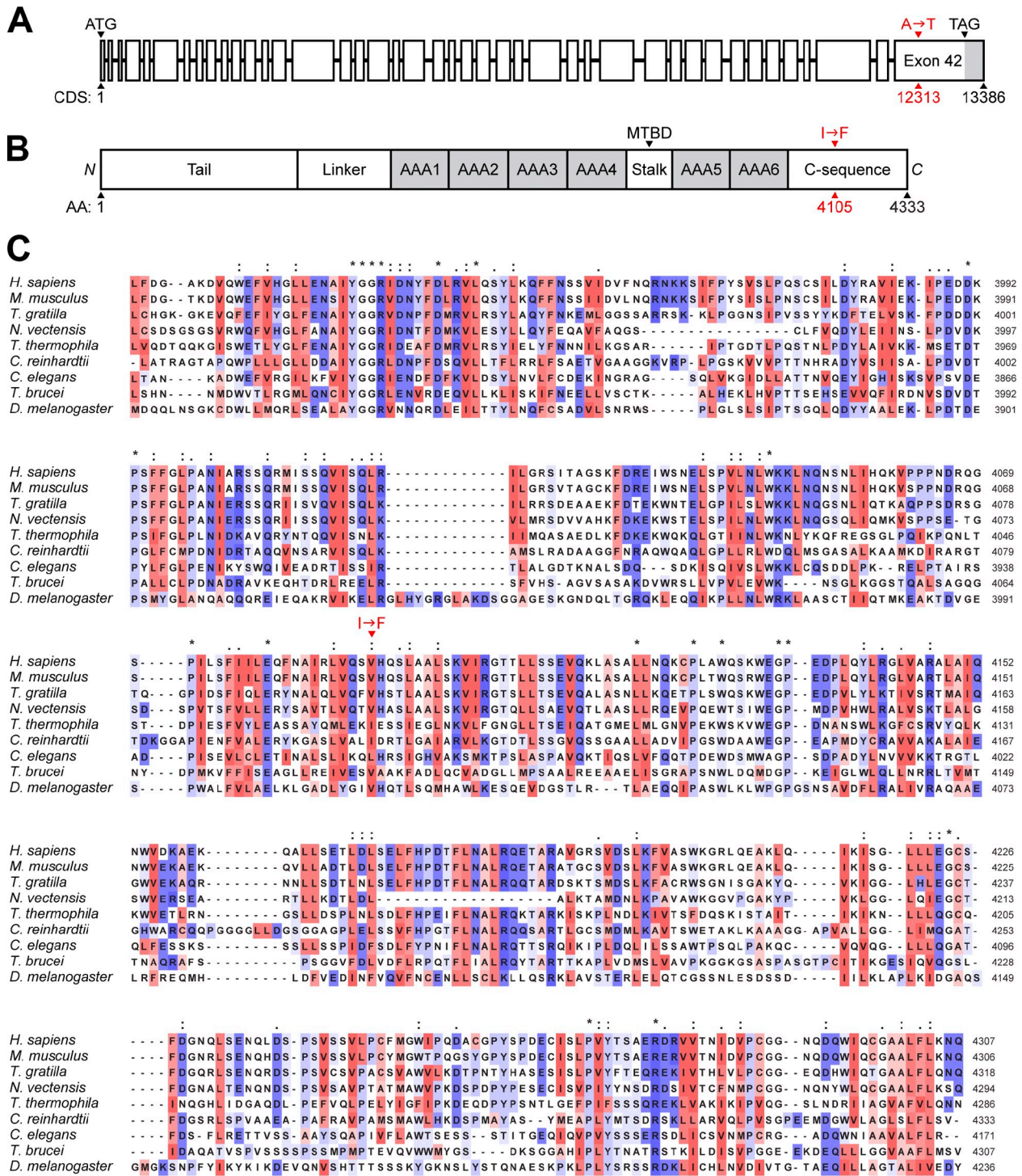


Figure S3. **The *dhc1b-3* mutation is a substitution of a single conserved residue in the C-sequence of DHC1b.** (A) Location of the *dhc1b-3* mutation in relation to the *Chlamydomonas* DHC1b gene, with boxes representing exons and the gray area signifying the 3' UTR. CDS, coding sequence. (B) Location of the amino acid substitution in relation to the DHC1b peptide. Characterized protein regions, such as the six ATPase domains, are labeled. AA, amino acid; MTBD, microtubule-binding domain. (C) ClustalX alignment of the DHC1b C-sequences (as defined by Numata et al., 2011) from human (*Hs*), mouse (*Mm*), sea urchin (*Tg*), cnidarian (*Nv*), *Tetrahymena* (*Tt*), *Chlamydomonas* (*Cr*), nematode (*Ce*), trypanosome (*Tb*), and fly (*Dm*). Standard Clustal conservation notation is shown ("*" for complete identity, ":" for strong group full conservation, and "." for weak group full conservation). The coloration represents conserved hydrophobic (red) and hydrophilic (blue) amino acids. The *dhc1b-3* mutation affects a hydrophobic residue with "strong group full conservation," located in the middle of a small conserved hydrophobic domain spanning 20–30 amino acids. Color values were taken from the Kyte-Doolittle hydrophobicity scale (Kyte and Doolittle, 1982), and whether a residue is conserved enough to be colored was decided by the ClustalX Zappo color assignment rules.

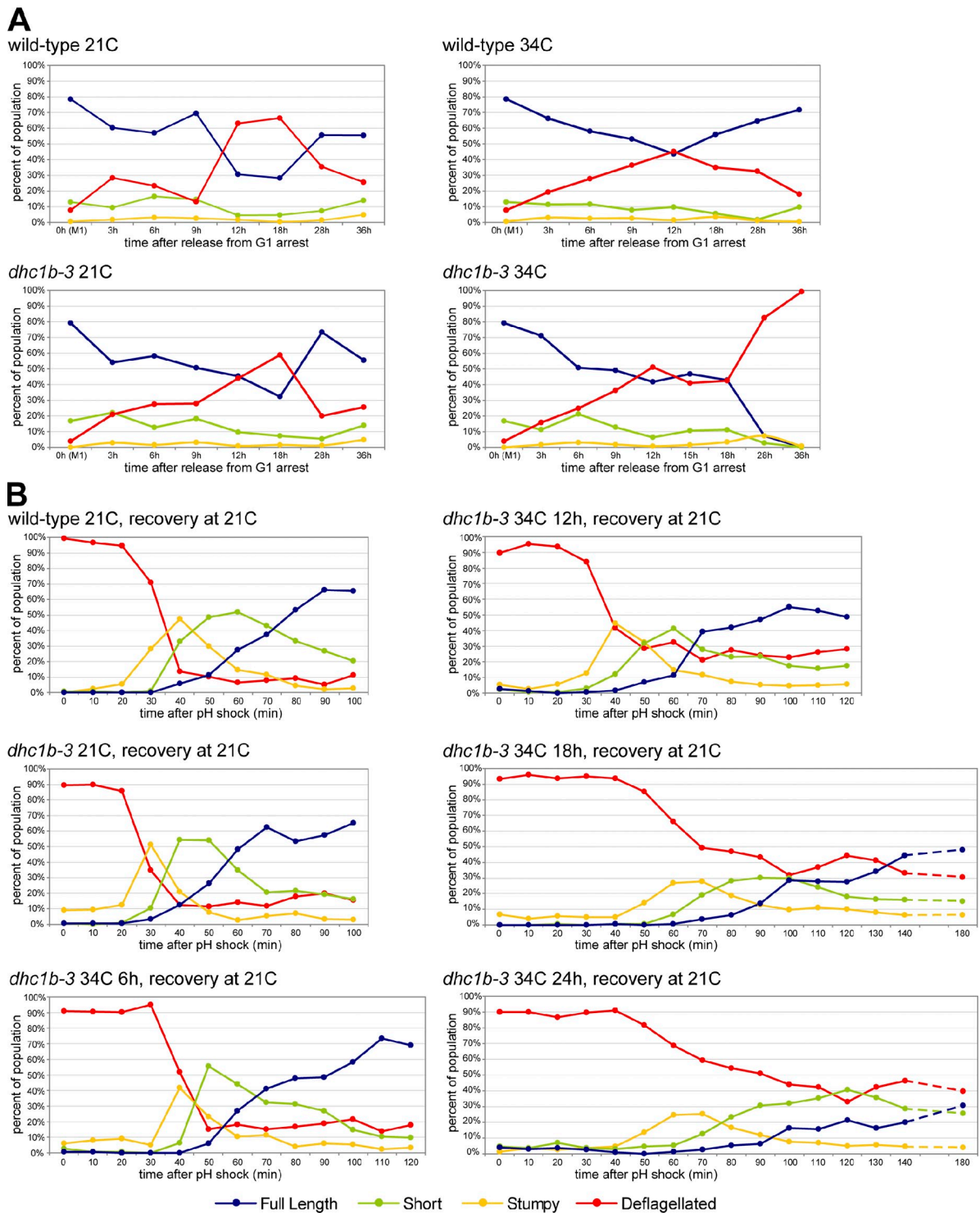


Figure S4. Kinetics of *dhc1b-3* flagellar loss in synchronized dividing cells and *dhc1b-3* flagellar regeneration at 21°C after preincubation at 34°C. In these experiments, flagellar length was judged visually and cells were binned into deflagellated, stumpy, short, and full length categories. (A) Log-phase cultures were synchronized by G1 growth arrest in M media in the dark for 12 h and then released from arrest by diluting cultures into TAP media in the light. At 21°C, both wild-type and *dhc1b-3* cultures went through a partially synchronized cell division at 12–18 h (~50% deflagellated and clumpy cells). After cell division, cells in both cultures regrew flagella. At 34°C, wild-type cells regrew flagella after cell division, but mutant cells were deflagellated. This implies that the *dhc1b-3* mutant can maintain old flagella at 21 and 34°C, but can only assemble new flagella at 21°C. (B) *dhc1b-3* cells were incubated for increasing periods of time (0, 6, 12, 18, and 24 h) at 34°C, deflagellated with pH shock, and then allowed to recover at 21°C. The kinetics of flagellar regeneration at 21°C were the same for wild-type and *dhc1b-3* cells that were not preincubated at 34°C. *dhc1b-3* cells that spent longer at 34°C before pH shock showed delayed onset of regeneration and slower kinetics of flagellar growth. $N_{\text{cells}} = 6,222$ (A), 14,907 (B) from single experiments.

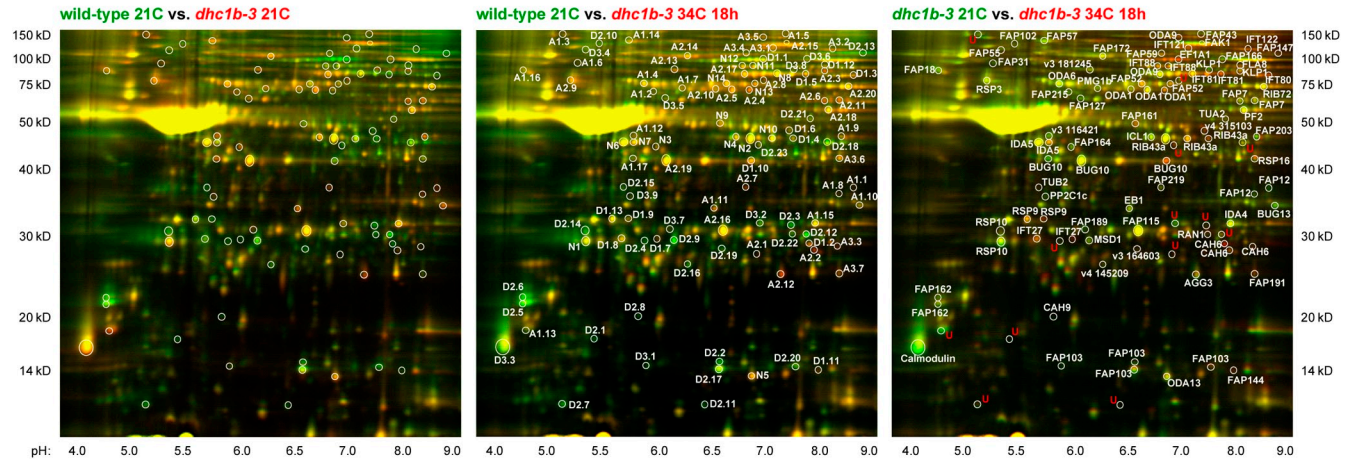
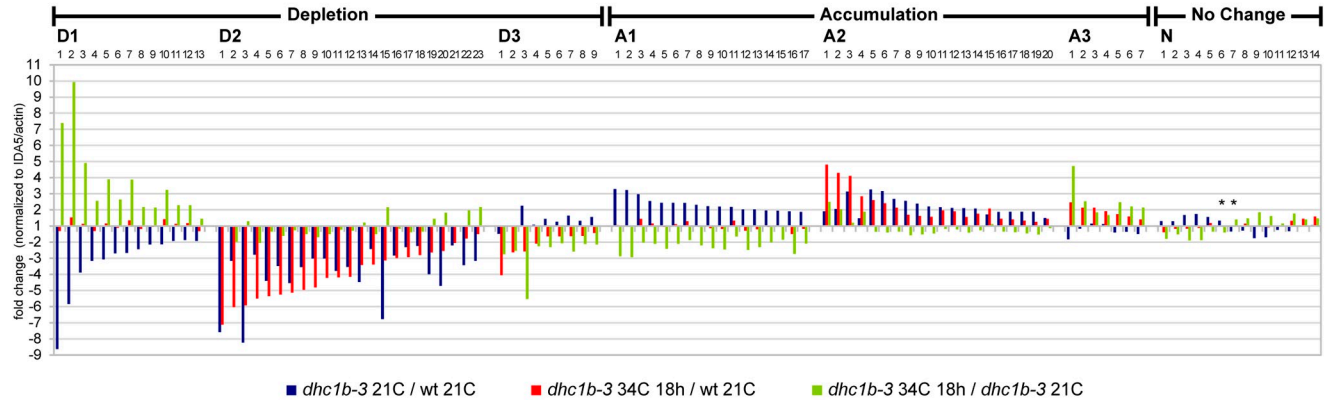
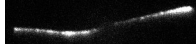
A**B**

Figure S5. **Complete results from the 2D-DIGE comparative proteomic analysis of isolated flagella (accompanies Table S3).** (A) Fluorescence images of the 2D-DIGE analytical gel, showing the abundance of proteins in the three samples relative to each other (wild-type 21°C, *dhc1b-3* 21°C, *dhc1b-3* 34°C 18 h; green and red spots = more protein in one sample than the other, yellow spots = equal abundance). The large yellow spot at ~50 kD is tubulin. White circles indicate spots that were picked for identification by mass spectrometry. In the center image, the spots are labeled with accumulation profile numbers that correspond to the graph in B and listings in Table S3. In the right image, the spots are labeled with their protein identities (U = unidentified). (B) Quantification of the fold change for every spot picked for mass spectrometry identification. Fold change is a spot's relative fluorescence between two samples on the analytical gel (1 = equal abundance for both samples). Spots were binned into classes based on whether they experienced accumulation or depletion in *dhc1b-3* flagella at 21°C, 34°C, or both vs. wild-type flagella at 21°C (D1 = depleted only at 21°C; D2 = depleted at 21 and 34°C; D3 = depleted only at 34°C; A1 = accumulated only at 21°C; A2 = accumulated at 21 and 34°C; A3 = accumulated only at 34°C). The two IDA5 actin spots in the "no change" class (marked with asterisks) have the smallest changes between the three samples and were used as controls to normalize the abundance of all of the other spots on the gel. Please see Table S3 for the identity and evaluation of each spot.



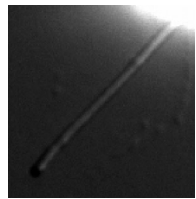
Video 1. **TIRF imaging of wild-type KAP-GFP flagella at 21°C.** Corresponds to the kymograph in Fig. 2 A (left). 30 s, real-time play speed. Images were acquired with objective-based TIRF at ~ 30 frames per second on a microscope (Eclipse Ti; Nikon).



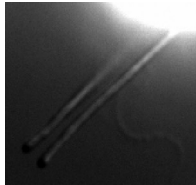
Video 2. **TIRF imaging of *dhc1b-3* KAP-GFP flagella at 21°C.** Corresponds to the kymograph in Fig. 2 A (middle). 34 s, real-time play speed. Images were acquired with objective-based TIRF at ~ 30 frames per second on a microscope (Eclipse Ti; Nikon).



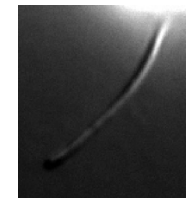
Video 3. **TIRF imaging of *dhc1b-3* KAP-GFP flagella at 34°C 12 h.** Corresponds to the kymograph in Fig. 2 A (right). 26 s, real-time play speed. Images were acquired with objective-based TIRF at ~ 30 frames per second on a microscope (Eclipse Ti; Nikon).



Video 4. **DIC imaging of wild-type flagellum at 21°C.** Corresponds to the kymograph in Fig. 2 B (left). 30 s, real-time play speed. Images were acquired at ~ 24 frames per second on a microscope (Eclipse Ti; Nikon) with DIC optics and a rotating stage.



Video 5. **DIC imaging of *dhc1b-3* flagellum at 21°C.** Corresponds to the kymograph in Fig. 2 B (middle). 32 s, real-time play speed. Images were acquired at ~ 24 frames per second on a microscope (Eclipse Ti; Nikon) with DIC optics and a rotating stage.



Video 6. **DIC imaging of *dhc1b-3* flagellum at 34°C 12 h.** Corresponds to the kymograph in Fig. 2 B (right). 29 s, real-time play speed. Images were acquired at ~ 24 frames per second on a microscope (Eclipse Ti; Nikon) with DIC optics and a rotating stage.

Table S1. Primers used to sequence the DHC1b cDNA, including UTRs

PCR fragment position	Forward primer	Reverse primer
-93-430 bp (5' UTR)	AAGCCCTGTGAACTTTGACG	CCAGCAGCTCAGACAAAACG
364-990 bp	CCGCTTATCAAGTCACAGACC	CCAGTCTCCACCAGAGC
973-1481 bp	GCTCTGGTGGAGGACTGG	AGGGTCTCCTTCTCACTCTGC
1399-2020 bp	GTGTTTGC GGAGATCAAGC	GTTTCAGCACCATACCGTACC
1866-2504 bp	GAACAGCCACGTGAAGACG	CGAAACACCATCTTGACTTCC
2405-3062 bp	GGGACTTCCAGCTGTACAAGG	ATGTCCTCCACGGTGTAGC
2875-3504 bp	CGCATCGAGTGCTACAAGG	TGGCAGTCAATCTTAAACG
3408-3982 bp	GGCCAAGATCCAGGAGTACG	TAAGCTCAGTCAGCGCCTTG
3863-4520 bp	TCACGCTCACGCAITTC	GCTTGGCCAGAAATCTCC
4363-5009 bp	GCTGATATCCCTGGCATCC	TCCTGGCCAGGACTCG
4868-5506 bp	AGCTGGTGACCGAGATGC	CCACCTCCACAGTCTTGTC
5427-6031 bp	CCAGCAGATTCAGACCATCC	GCGTCAGTCCATCTTGC
5893-6569 bp	CTCACCTTTGACGACAACCTCC	GTGGCTGCACCTTGAGC
6383-7061 bp	ACCGCCTGCTGACTATGC	ACACCATGATGCGCTTG
6924-7593 bp	GGTACACCCGAGGTGACG	GTAGTGCGGGTAGTCGTTGG
7411-8095 bp	ATCGTACCATGGGCTACC	CCATGTTGTGCATGTACGC
7897-8450 bp	GAGCTAGTGGCGGAAAAGC	CGGAACATCTCGTTTGAGG
8369-9055 bp	ACTCGTACGCCTTCTCAGC	TCTCAGCGATGTTATCCTTCTTG
8934-9511 bp	CGAGGAGGTGGAGATGAAGG	CCTCGCTGGTCTTCTTGC
9398-10001 bp	AGAGCAGTCTGGAGGAGAGC	TCCAGTGTGGTGGTGAAGC
9859-10539 bp	ATCCTCAACTCCACCTCTCG	GGGTCGGTACACCTCACG
10436-11057 bp	AGCTGAACGAGACCAAGACG	AGCTGCTGGAAGGGGTTTC
10822-11377 bp	CAGTGGGCCTTCTTCTCTG	TCTTGAGCACCAGCCAGTC
10822-11432 bp	CAGTGGGCCTTCTTCTCTG	AGCATCAGCAGCTCCTTCTC
10829-11452 bp	CCTTCTTCTGGGCAAGC	GGAAGTTGTCGTGCTTCTGC
11363-11987 bp	GGCTGGTGCTCAAGAACG	GCGGAGATGATGGACACG
11845-12699 bp (spans mutation)	CAAGTGCTGCTGACCTTCC	CITGAGCTTGGCGGTTTC
12399-12968 bp	GCTGGCGGACGTTATTCC	ACCCAGCCATCCATCTCC
12835-13349 bp (3' UTR)	TGGCTGCACAAGGACTCG	TGTCAGAAATCAAACATCAGTCG

Primer sequences are listed from 5' to 3'.

Table S2. Quantification of the Western blots shown in Fig. 5 B, normalized by the levels of α -tubulin

Flagellar protein	Wild-type 21°C	Wild-type 34°C 18 h	<i>dhc1b-3</i> 21°C	<i>dhc1b-3</i> 34°C 6 h	<i>dhc1b-3</i> 34°C 12 h	<i>dhc1b-3</i> 34°C 18 h
DHC1b	1.0	1.1	-3.6	-4.8	-11.8	-28.2
D1bLIC	1.0	1.1	-2.2	-8.5	-4.9	-8.0
FLA10	1.0	1.2	1.6	1.7	1.3	1.7
KAP	1.0	1.6	1.9	2.1	2.1	1.6
IFT27	1.0	1.1	1.4	2.8	2.9	2.8
IFT81	1.0	-3.5	-3.2	1.6	1.6	3.8
IFT172	1.0	-1.4	1.8	1.1	1.7	2.4
IFT122	1.0	1.3	2.1	2.1	2.5	2.4
IFT140	1.0	-3.1	1.6	3.0	4.0	4.1
BBS4	1.0	-1.2	-1.4	-1.2	1.1	1.4
ODA-IC2	1.0	-5.3	2.2	2.6	2.7	2.5
Rib43a	1.0	1.4	1.2	-1.2	1.3	1.0

Numbers indicate the fold increase or fold decrease in protein abundance relative to wild-type flagella at 21°C. It should be noted that the immunoblots were visualized with film, which cannot be quantified with high precision due to saturation of the signal. Changes greater than twofold are bold to indicate that they are likely significant.

Table S3. Full descriptions of proteins identified by 2D-DIGE, clustered by accumulation profile (accompanies Fig. S5).

D1: Identification

Spot	Protein	Counts	MW	PI	Spot shift
D1.1	eEF1A1 (v4 134382) and/or TUA1	17, 12	100	7.0	
D1.2	CAH6 (v4 133249), carbonic anhydrase 6	12	30	7.9	-0.5 pH, +0.5 kD
D1.3	IFT81 or IFT80 (FAP167), IFT-B	16, 11	84	8.7	
D1.4	Rib43a , protofilament ribbons of flagellar microtubules	17	48	7.7	
D1.5	IFT81 , IFT-B	34	86	7.9	
D1.6	v4 315103 (v3 190810, estExt_fgenesh2_pg.C_200113) similar to HSP70	15	50	7.5	
D1.7	IFT27 (FAP156, RabL4, small Rab-related GTPase), IFT-B	5	30	6.0	
D1.8	IFT27 (FAP156, RabL4, small Rab-related GTPase), IFT-B	8	30	5.6	-0.4 pH
D1.9	PF17 , radial spoke protein 9 (RSP9)	12	34	5.8	
D1.10	BUG10 (FAP252)	15	42	6.9	
D1.11	FAP144	7	15	8.1	
D1.12	FLA8 , kinesin-2 anterograde motor or KLP1	5, 8	95	8.0	
D1.13	PF17 , radial spoke protein 9 (RSP9)	10	34	5.6	-0.2 pH

D1: Validation

Spot	Predicted MW	Predicted PI	Flagellar proteome? (counts, fraction)	Notes
D1.1	51, 50	8.7, 5.0	Yes (28, MM , KCI , T, Ax), Yes (31, Ax , KCI , MM , T)	EF1A1-tubulin dimer? Cargo?
D1.2	28	7.0	Yes (7, T , MM , KCI)	Anterograde cargo?
D1.3	77, 84	6.3, 6.6	Yes (26, MM), Yes (7, MM)	IFT-B
D1.4	43	6.0	Yes (33, Ax , T)	Anterograde cargo? Unmodified?
D1.5	77	6.3	Yes (26, MM)	IFT-B, unmodified?
D1.6	44	6.0	Yes (6, Ax)	Anterograde cargo?
D1.7	23	5.3	Yes (8, MM)	IFT-B, GDP bound?
D1.8	23	5.3	Yes (8, MM)	IFT-B, GTP bound?
D1.9	30	5.0	Yes (17, T , Ax , KCI, MM)	Anterograde cargo?
D1.10	39	5.7	Yes (29, T , Ax , KCI, MM)	Anterograde cargo? Unmodified?
D1.11	16	7.0	Yes (9, Ax , T)	Anterograde cargo?
D1.12	86, 83	6.6, 6.5	Yes (19, MM , KCI , T), Yes (31, KCI , MM)	Anterograde IFT motor
D1.13	30	5.0	Yes (17, T , Ax , KCI, MM)	-0.4 pH spot has "no change" profile

D2: Identification

Spot	Protein	Counts	MW	PI	Spot shift
D2.1	Unidentified	-	17	5.4	
D2.2	FAP103 or v3 180172 (fgenes2_pg.C_scaffold_8100021)	5, 3	16	6.6	-1.0 pH, +0.5 kD
D2.3	Unidentified	-	33	7.4	
D2.4	v3 184650 (estExt_fgenes2_kg.C_300011)	5	30	5.9	
D2.5	FAP162	12	22	4.7	
D2.6	FAP162 or SKP1 (v4 128004), E3 ubiquitin ligase	8, 2	23	4.7	+1 kD
D2.7	unidentified	-	10	5.2	
D2.8	CAH9 (v3 196834), carbonic anhydrase 9	2	21	5.8	
D2.9	MSD1 (v4 53941), superoxide dismutase [Mn], mitochondrial	8	30	6.2	
D2.10	FAP102 or TUA2 , alpha tubulin 2	5, 12	140	5.5	
D2.11	FKB12 , peptidyl-prolyl cis-trans isomerase, FKBP-type	4	10	6.5	
D2.12	CYB5_3 (e_gwH.2.411.1)	5	31	7.9	
D2.13	FAP147 , similar to myc-binding	16	105	8.9	
D2.14	RSP10 , radial spoke protein 10	6	32	5.4	+2 kD
D2.15	TUB2 , beta tubulin 2	11	38	5.7	
D2.16	v4 145209 , Chlre2_kg.scaffold_13000122, adenine phosphoribosyltransferase	6	27	6.3	
D2.17	FAP103 or v3 180172 (fgenes2_pg.C_scaffold_8100021)	5	15	6.6	-1.0 pH, -0.5 kD
D2.18	IDH3 (v4 196567), isocitrate dehydrogenase, NADP-dependent, mitochondrial	18	47	8.2	
D2.19	v3 164603 (fgenes2_kg.C_scaffold_45000028) glutathione S-transferase	4	29	6.7	
D2.20	FAP103 or v3 180172 (fgenes2_pg.C_scaffold_8100021)	5	15.5	7.6	
D2.21	TUA2 or TUB2 or SHMT2 (v4 196354), serine hydroxymethyltransferase 2	10, 14, 10	58	8.0	
D2.22	RAN1 , Ran-like small GTPase	3	32	7.6	
D2.23	Unidentified	-	48	6.9	

D2: Validation

Spot	Predicted MW	Predicted PI	Flagellar proteome? (counts, fraction)	Notes
D2.1	-	-	-	
D2.2	17, 16	6.2, 6.2	Yes (9, MM), No	Glutathiolation target
D2.3	-	-	-	
D2.4	22	6.5	-	
D2.5	19	4.5	Yes (8, MM)	
D2.6	19, 18	4.5, 4.6	Yes (8, MM), No	
D2.7	-	-	-	
D2.8	13	6.4	?	
D2.9	24	5.9	No?	Contaminant? Flagellar redox?
D2.10	82, 50	8.7, 5.0	Yes (13, Ax, KCI), Yes (35, Ax, KCI, T, MM)	Redox-sensitive
D2.11	12	6.0	-	
D2.12	12	9.0	-	
D2.13	97	6.7	Yes (9, KCI)	
D2.14	24	5.0	Yes (9, T, Ax)	Higher MW than "no change" spot
D2.15	50	4.8	Yes (27, KCI, MM, Ax, T)	MW too low for tubulin
D2.16	20	6.0	Yes (1, MM)	
D2.17	17, 16	6.2, 6.2	Yes (9, MM), No	Nucleoside diphosphate kinase
D2.18	54	9.0	No?	Contaminant? flagellar redox?
D2.19	24	5.5	?	Cannot verify ID in genome search
D2.20	17, 16	6.2, 6.2	Yes (9, MM), No	Unmodified?
D2.21	50, 50, 52	5.0, 4.8, 6.3	Yes, Yes (35, Ax, KCI, T, MM), ?	
D2.22	25	6.2	Yes (3, MM)	
D2.23	-	-	-	

D3: Identification

Spot	Protein	Counts	MW	PI	Spot shift
D3.1	FAP103 , nucleoside diphosphate kinase-like protein	3	15.5	5.8	-1.8 pH
D3.2	Unidentified	-	33	6.9	
D3.3	CAM1 , calmodulin	8	18	4.3	
D3.4	FAP55 or TUB2 , beta tubulin 2	22, 18	115	5.4	
D3.5	FAP127	11	70	6.1	
D3.6	FAP166	9	32	6.1	
D3.7	FAP189 or FAP58	12, 11	100	7.8	
D3.8	KLP1 , kinesin-like protein 1	22	90	7.7	-0.5 pH
D3.9	PP2C1c , protein phosphatase 2C catalytic subunit isoform 1	15	37	5.8	

D3: Validation

Spot	Predicted MW	Predicted PI	Flagellar proteome? (counts, fraction)	Notes
D3.1	17	6.2	Yes (9, MM)	Gutthiolation target
D3.2	-	-	-	
D3.3	18	4.3	Yes (9, MM , Ax , KCI , T)	
D3.4	112, 50	5.1, 4.8	Yes (27, Ax , T), Yes (27, KCI , MM , Ax , T)	
D3.5	51	5.3	Yes (11, Ax , T)	
D3.6	24	5.5	Yes (7, Ax , T)	
D3.7	102, 101	6.8, 6.6	Yes (6, Ax , KCI), Yes (26, Ax , T, KCI)	
D3.8	83	6.5	Yes (31, KCI , MM)	
D3.9	29	5.6	Yes? (14, T)	

A1: Identification

Spot	Protein	Counts	MW	PI	Spot Shift
A1.1	FAP12 , lipase domain	7	38	8.7	
A1.2	FAP215 , conserved	8	73	6.0	
A1.3	Unidentified	-	200	5.1	
A1.4	ODA6 , flagellar outer dynein arm intermediate chain 2 (ODA-IC2)	12	80	5.9	
A1.5	FAP43 , conserved	16	200	7.3	
A1.6	FAP31 , conserved TPR repeat, or FAP197	18, 6	95	5.4	
A1.7	PGM1b (v4 161085), phosphoglycerate mutase, or PGM1a or ODA1	13, 13, 11	75	6.3	
A1.8	FAP12 , lipase domain	5	37	8.4	-0.3 pH, -1 kD
A1.9	FAP203 , weakly similar to HSP70	8	48	8.4	
A1.10	BUG13 (FAP90), basal body protein of unknown function	10	35	8.9	
A1.11	EB1	12	35	6.6	
A1.12	v3 116421 (e_gwW.1.241.1)	13	49	5.8	
A1.13	Unidentified	-	18	4.7	
A1.14	FAP57 , conserved	10	150	5.8	
A1.15	IDA4 , flagellar inner arm dynein light chain p28	18	33	8.0	
A1.16	FAP18 , high E/A/P content protein	15	90	4.7	
A1.17	BUG10 (FAP252)	19	44	5.8	-1.1 pH

A1: Validation

Spot	Predicted MW	Predicted PI	Flagellar proteome? (counts, fraction)	Notes
A1.1	54	5.7	Yes (25, T , MM , KCI , Ax)	MW too low for FAP12
A1.2	52	5.0	Yes (6, MM)	
A1.3	-	-	-	
A1.4	63	5.4	Yes (9, T)	
A1.5	178	8.5	Yes (54, KCI , Ax , T)	
A1.6	74, 33	6.5, 6.0	Yes (9, MM), Yes (6, KCI)	
A1.7	61, 67, 62	5.6, 5.6, 5.8	Yes (6, MM), ?, Yes (18, T)	
A1.8	54	5.7	Yes (25, T , MM , KCI , Ax)	MW too low for FAP12
A1.9	19	7.2	Yes (6, Ax , KCI, MM)	MW too high for FAP203
A1.10	28	8.6	Yes (15, T , Ax , KCI)	
A1.11	31	5.7	Yes (2, MM)	
A1.12	31	5.3	?	Cannot verify ID in genome search
A1.13	-	-	-	
A1.14	120	6.2	Yes (26, Ax , KCI , T)	
A1.15	29	6.7	Yes (21, KCI , T , Ax , MM)	
A1.16	55	4.4	Yes (17, KCI , MM , T , Ax)	
A1.17	39	5.7	Yes (29, Ax , T , KCI, MM)	

A2: Identification

Spot	Protein	Counts	MW	PI	Spot shift
A2.1	Unidentified	-	28	6.8	
A2.2	CAH6 (v4 133249), carbonic anhydrase 6	9	29	8.0	-0.4 pH, -0.5 kD
A2.3	KLP1 , kinesin-like protein 1	35	90	8.2	
A2.4	ODA1 , flagellar outer dynein arm-docking complex protein 2 (ODA-DC2)	35	75	6.9	
A2.5	ODA1 , flagellar outer dynein arm-docking complex protein 2 (ODA-DC2)	23	75	6.7	-0.2 pH
A2.6	FAP7	21	68	8.2	-0.3 pH
A2.7	FAP219	11	38	6.9	
A2.8	Unidentified	-	80	7.1	
A2.9	PF14 , radial spoke protein 3 (RSP3)	20	80	5.3	
A2.10	ODA1 , flagellar outer dynein arm-docking complex protein 2 (ODA-DC2)	17	75	6.5	-0.4 pH
A2.11	FAP7	21	68	8.5	
A2.12	AGG3 (FAP142), flavodoxin similar to quinone reductase	6	26	7.3	
A2.13	v3 181245 (fgenes2_pg.C_scaffold_721000001)	15	90	6.2	
A2.14	FAP172 or Dhc1b	24, 33	105	6.3	
A2.15	FAK1 , ODA5-assoc. flag. adenylate kinase Or FAK2 (v3 196915) or FAP66	7, 7, 18	145	7.5	
A2.16	FAP115	8	32	6.7	
A2.17	ODA9 (ODA-IC1) Or v3 153411 (Chlre2_kg.scaffold_62000038)	28	85	6.8	
A2.18	PF2 , component 4 of the dynein regulatory complex (DRC4)	9	62	8.2	
A2.19	BUG10 (FAP252)	20	44	6.1	-0.8 pH
A2.20	Rib72 , pf ribbons of flag MT, nucleoside-diphosphokinase regulatory subunit p72	22	77	8.6	

A2: Validation

Spot	Predicted MW	Predicted PI	Flagellar proteome? (counts, fraction)	Notes
A2.1	-	-	-	
A2.2	28	7.0	Yes (7, T , MM , KCl)	
A2.3	83	6.5	Yes (31, KCl , MM)	Unmodified?
A2.4	62	5.8	Yes (18, T)	
A2.5	62	5.8	Yes (18, T)	
A2.6	55	6.7	Yes (20, Ax , T , KCl , MM)	
A2.7	21	9.7	Yes (5, Ax , T)	
A2.8	-	-	-	
A2.9	57	4.9	Yes (27, Ax , T , KCl)	
A2.10	62	5.8	Yes (18, T)	
A2.11	55	6.7	Yes (20, Ax , T , KCl , MM)	
A2.12	21	6.3	Yes (9, MM)	
A2.13	72	11.7	?	
A2.14	64, 481	5.5, 6.1	Yes (7, Ax , T), Yes (88, MM , KCl, Ax)	
A2.15	71, 54, 146	6.6, 5.9, 6.0	Yes (25, T , KCl , MM , Ax), Yes (23, MM , KCl, T)	
A2.16	27	5.8	Yes (11, Ax , T , KCl, MM)	
A2.17	76, 30	5.9, 10.1	Yes (14, T), -	
A2.18	55	8.4	Yes (16, Ax , T , KCl)	
A2.19	39	5.7	Yes (29, Ax , T , KCl, MM)	
A2.20	72	7.2	Yes (44, Ax , T , KCl, MM)	

A3: Identification

Spot	Protein	Counts	MW	PI	Spot Shift
A3.1	IFT121 (IFT122B, IFTA-1, FAP118 / FBB1), cargo adaptor, IFT-A	25	120	7.2	
A3.2	IFT122 (IFT122A, DAF-10, FAP80) or v4 299962 , subtilisin-like serine protease	11, 8	120	8.2	
A3.4	CAH6 (v4 133249), carbonic anhydrase 6	8	29.5	8.4	
A3.4	FAP59 , conserved	10	105	6.8	
A3.5	ODA9 , flagellar outer dynein arm intermediate chain 1 (ODA-IC1)	6	200	7.0	+115 kD
A3.6	RSP16 , radial spoke protein 16	16	44	8.4	
A3.7	FAP191 , similar to quinone reductase or AGG3	1, 5	26	8.4	

A3: Validation

Spot	Predicted MW	Predicted PI	Flagellar Proteome? (counts, fraction)	Notes
A3.1	136, 132	6.0, 6.3	Yes (11, MM , KCl)	IFT-A
A3.2	137, 85	6.6, 9.0	Yes (16, MM , KCl, T), Yes (2, Ax)	IFT-A
A3.4	28	7.0	Yes (7, T , MM , KCl)	Retrograde cargo?
A3.4	101	6.0	Yes (26, Ax , T , KCl)	Retrograde cargo?
A3.5	76	5.9	Yes (14, T)	Retrograde cargo? MW too high.
A3.6	39	6.7	Yes (26, T , Ax , KCl, MM)	Retrograde cargo?
A3.7	28, 21	7.1, 6.3	Yes (6, MM), Yes (9, MM)	Same MW as AGG3 (+1.1 pH), cargo?

No Change: Identification

Spot	Protein	Counts	MW	PI	Spot shift
N.1	RSP10 , radial spoke protein 10	6	30	5.4	
N.2	Rib43a , protofilament ribbons of flagellar microtubules	24	48	6.9	-0.8 pH
N.3	FAP164	12	47	6.0	
N.4	ICL1 , isocitrate lyase <i>or</i> Rib43a	14, 12	48	6.7	
N.5	ODA13 , flagellar outer dynein arm light chain 6 (ODA-LC6)	6	14	6.9	
N.6	IDA5 , actin	21	48	5.7	-0.1 pH
N.7	IDA5 , actin	16	48	5.8	
N.8	IFT81 , IFT-B	21	86	7.3	-0.6 pH
N.9	FAP161	12	54	6.6	
N.10	Rib43a , protofilament ribbons of flagellar microtubules	23	48	7.3	-0.4 pH
N.11	IFT88 , IFT-B or METE , Vit B12 (cobalamin) independent methionine synthase	20, 16	94	7.0	
N.12	IFT88 , IFT-B	15	94	6.8	-0.2 pH
N.13	FAP52	19	80	7.0	
N.14	FAP52	30	80	6.7	-0.3 pH

No Change: Validation

Spot	Predicted MW	Predicted PI	Flagellar Proteome? (counts, fraction)	Notes
N.1	24	5.0	Yes (9, T , Ax)	− 2 kD from “D2” spot
N.2	43	6.0	Yes (33, Ax , T)	
N.3	48	5.9	Yes (8, MM)	
N.4	46, 43	5.9, 6.0	Yes (3, MM), Yes (33, Ax , T)	
N.5	14	6.3	Yes (6, T)	
N.6	42	5.3	Yes (22, KCI , Ax , T , MM)	Normalization control
N.7	42	5.3	Yes (22, KCI , Ax , T , MM)	Normalization control
N.8	77	6.3	Yes (26, MM)	IFT-B, lower pH than “D1” spot
N.9	43	5.6	Yes (8, Ax , KCI)	
N.10	43	6.0	Yes (33, Ax , T)	
N.11	86, 87	5.9, 6.0	Yes (14, MM)	IFT-B
N.12	86	5.9	Yes (14, MM)	IFT-B
N.13	67	5.8	Yes (30, Ax , T , KCI)	
N.14	67	5.8	Yes (30, Ax , T , KCI)	

Color key for columns: “Protein” red = unidentified or low confidence hit; “Predicted MW” and “Predicted PI” blue = good match (<20 kD, <1 pH difference), red = bad match; “Flagellar Proteome?” bold = major fraction(s), green = major fraction is in the membrane + matrix, red = not found in proteome (Pazour et al., 2005). Proteome fractions are listed in order of abundance. “Spot shift” is the estimated distance on the 2D analytical gel from an additional spot with the same ID.

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