bop5 mutations reveal new roles for the IC138 phosphoprotein in the regulation of flagellar motility and asymmetric waveforms

Kristyn E. VanderWaal^a, Ryosuke Yamamoto^b, Ken-ichi Wakabayashi^c, Laura Fox^b, Ritsu Kamiya^c, Susan K. Dutcher^d, Phillip V. Bayly^e, Winfield S. Sale^b, and Mary E. Porter^a

^aDepartment of Genetics, Cell Biology, and Development, University of Minnesota, Minneapolis, MN 55455; ^bDepartment of Cell Biology, Emory University School of Medicine, Atlanta, GA 30322; ^cDepartment of Biological Sciences, Graduate School of Science, University of Tokyo, Tokyo 113-0033, Japan; ^dDepartments of Genetics and Cell Biology and Physiology, Washington University in St. Louis, St. Louis, MO 63110; ^eDepartment of Mechanical, Aerospace and Structural Engineering, Washington University in St. Louis, St. Louis, MO 63130

ABSTRACT I1 dynein, or dynein f, is a highly conserved inner arm isoform that plays a key role in the regulation of flagellar motility. To understand how the IC138 IC/LC subcomplex modulates I1 activity, we characterized the molecular lesions and motility phenotypes of several bop5 alleles. bop5-3, bop5-4, and bop5-5 are null alleles, whereas bop5-6 is an intron mutation that reduces IC138 expression. I1 dynein assembles into the axoneme, but the IC138 IC/LC subcomplex is missing. bop5 strains, like other I1 mutants, swim forward with reduced swimming velocities and display an impaired reversal response during photoshock. Unlike mutants lacking the entire I1 dynein, however, bop5 strains exhibit normal phototaxis. bop5 defects are rescued by transformation with the wild-type IC138 gene. Analysis of flagellar waveforms reveals that loss of the IC138 subcomplex reduces shear amplitude, sliding velocities, and the speed of bend propagation in vivo, consistent with the reduction in microtubule sliding velocities observed in vitro. The results indicate that the IC138 IC/LC subcomplex is necessary to generate an efficient waveform for optimal motility, but it is not essential for phototaxis. These findings have significant implications for the mechanisms by which IC/ LC complexes regulate dynein motor activity independent of effects on cargo binding or complex stability.

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INTRODUCTION

The axonemal dyneins are microtubule motors necessary for production of the elegant and sophisticated movement of cilia and flagella. Axonemal dyneins consist of two main classes: two- or threeheaded outer arm dyneins, which are responsible for generation of power and flagellar beat frequency (Brokaw and Kamiya, 1987), and multiple inner arm dyneins, which control the size and shape of the

flagella bends (Porter and Sale, 2000; Smith and Yang, 2004; King and Kamiya, 2009). The inner dynein arms consist of more than seven different subspecies that are localized to specific regions within each 96 nm repeat (Brokaw and Kamiya, 1987; Mastronarde et al., 1992; Brokaw, 1994; Porter et al., 1996; Porter and Sale, 2000; Kamiya, 2002; Nicastro et al., 2006; Wirschell et al., 2007; Bui et al., 2008; Yagi et al. 2009). Each inner arm dynein likely plays a unique role in the formation of the flagellar bend (Kamiya, 2002; Brokaw, 1994, 2009), but the specific function of each isoform is still unclear.

To begin to understand each inner arm dynein's role in motility, we have focused on one subspecies, known as I1 or f dynein (reviewed in Porter and Sale, 2000; Kamiya, 2002; Wirschell et al., 2007). I1 dynein is the most highly conserved and widely distributed inner arm dynein (Wickstead and Gull, 2007). The I1 complex is the only two-headed inner arm dynein, and it assembles as a trilobed structure near the base of radial spoke 1 (Figure 1). The two heads contain the motor domains of the two DHCs (dynein heavy chains),

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Abbreviations used: BOP, bypass of paralysis; DHC, dynein heavy chain; FAP, flagellar associated protein; HC, heavy chain; IC, intermediate chain; IDA, inner dynein arm; LC, light chain; MBO, move backward only; MIA, modifier of inner arms; PF, paralyzed flagellar; TAP, Tris acetate phosphate.

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FIGURE 1: Diagrammatic representation of 11 dynein substructure in the axoneme. This diagram illustrates the structure of the 11 dynein and the proposed interactions between 11 subunits based on previously published work (Myster et al., 1999; Perrone et al., 2000; Porter and Sale, 2000; Ikeda et al., 2009; Wirschell et al., 2009; modified from Bower et al., 2009).

1α and 1β, and the base consists of the DHC tail domains, intermediate chains (IC97, IC138, and IC140), light chains (LC7a, LC7b, LC8, TcTex1, and Tctex2b), and FAP120. (Goodenough and Heuser, 1985a, 1985b; Piperno *et al.*, 1990; Smith and Sale, 1991, 1992; Porter *et al.*, 1992; Myster *et al.*, 1997, 1999; Harrison *et al.*, 1998; Perrone *et al.*, 1998, 2000; Yang and Sale, 1998; DiBella *et al.*, 2004a, 2004b; Hendrickson *et al.*, 2004; Bower *et al.*, 2009; Ikeda *et al.*, 2009; Wirschell *et al.*, 2009). A collection of mutations in the genes encoding different I1 subunits, which results in the assembly of partial I1 dynein complexes in the axoneme, has been useful in determining the locations of and interactions between different subunits of the complex. Likewise, these mutant strains have given us insights into the function of each subunit (see Table 1).

The first I1 mutant to be extensively studied was pf9-3, which affects the DHC1 gene and is null for the 1α DHC subunit (Myster et al., 1997). Loss of the 1 DHC prevents the I1 complex from being assembled in the axoneme (Myster et al., 1997). Most mutations in the 1β DHC subunit (*ida2*) and IC140 (*ida7*) also block assembly of the I1 dynein (Perrone et al., 1998, 2000; Yagi et al., 2009). These mutant strains exhibit slow, forward-swimming velocities, reduced microtubule sliding, an inability to phototax, and abnormal flagellar waveforms (Perrone et al., 2000). Notably, transformation of the mutants with gene fragments sometimes results in the partial assembly of I1 dynein in the axoneme. For example, transformation with constructs encoding the N-terminal portions of the DHCs resulted in strains (pf9::DHC1 and ida2::DHC10) that assemble all I1 subunits except the 1α or 1β motor domain and display increased swim speeds and phototaxis ability (Myster et al., 1999; Perrone et al., 2000). More recent studies have revealed the distinct functional capability of each motor domain in the I1 dynein complex (Toba et al., 2011).

Another subunit thought to play a key role in the function of I1 dynein and control of flagellar bending is IC138. For example, in vitro studies revealed that changes in phosphorylation of IC138 correlate with changes in microtubule sliding velocities (Habermacher and Sale, 1997). In addition, analyses of the *mia* mutants, which cannot phototax and swim slowly, showed hyperphosphorylation of IC138 in the axoneme (King and Dutcher, 1997). These results suggested that IC138 is a regulatory phosphoprotein important for control of motility. The first *IC138* mutation, *bop5-1*, was isolated as a suppressor that could restore partial motility to the paralyzed

flagellar (pf) mutant pf10 (Dutcher et al., 1988). The DIC4/BOP5 gene encodes IC138¹, and the bop5-1 mutation produces a truncated protein with reduced activity, which supports the idea that IC138 is a key regulatory subunit (Hendrickson et al., 2004). When the full-length, wild-type DIC4 gene is transformed into bop5-1, the motility and protein defects are rescued (Hendrickson et al., 2004). Because bop5-1 is not a true null, however, further analysis of IC138 function was limited.

The first IC138 null to be characterized in detail was the *bop5-2* strain (Bower *et al.*, 2009). Analysis of *bop5-2* showed that IC138 is required for the assembly of four subunits at the base of the I1 dynein: IC138, LC7b, IC97, and FAP120, which is named the IC138 subcomplex (Figure 1 and Bower *et al.*, 2009). This study also showed that regulation of microtubule sliding velocities by the central pair/radial spoke phosphorylation pathway depends on the presence of the IC138 subcomplex (Bower *et al.*, 2009; Wirschell *et al.*, 2009). Similar to *bop5-1*, microtubule sliding defects in *bop5-2* and *bop5-2 pf17* were restored upon transformation with the wild-type *IC138* gene and reassembly of the IC138 subcomplex. Because *bop5-2* has additional mutations, however, it was not possible to discern how the loss of IC138 affects flagellar motility and cell behavior (Bower *et al.*, 2009).

In this study, we used the entire complement of bop5 mutants affecting IC138 (bop5-1 through bop5-6 [Hendrickson et al., 2004; Bower et al., 2009; Ikeda et al., 2009]) to determine the specific roles of IC138 in regulating flagellar motility. On the basis of earlier studies of I1 dynein mutants, we postulated that assembly of IC138 would be required to control swimming speed, flagellar waveform, and phototaxis. We determined that, although the I1 dynein is necessary for normal phototaxis, the IC138 subcomplex is not. In contrast, the loss of any subunit of the I1 complex is sufficient to impair cell reversal during the photoshock response. Also, we characterized the flagellar waveform of the bop5-3 mutant and compared it to those previously described for wild type and ida3 (missing the entire I1) strains (Bayly et al., 2010) to determine how waveform might be affected by loss of the IC138 subcomplex. We observed that the bop5-3 waveform is intermediate between wild type and ida3. Mutations in I1 do not affect the maximum and minimum curvature of the waveform, but they do affect the sliding velocity and shear amplitude of the wave patterns during forward swimming. Thus, we propose that the IC138 IC/LC subcomplex is critical for control of microtubule sliding and flagellar waveform, but it is not required for phototaxis. Given the high degree of I1 dynein subunit conservation, these observations have important implications for the mechanisms by which IC/LC subcomplexes regulate ciliary and flagellar motility in many species.

RESULTS

Molecular characterization of the mutations in bop5-3, bop5-4, bop5-5, and bop5-6

To determine the nature of the *bop5* mutations that lead to decreased *IC138* expression in the collection first described in Ikeda *et al.* (2009), the entire *IC138* gene in each strain was recovered by PCR and sequenced directly (Figure 2). In *bop5-3*, we were unable to amplify a small region at the 5' end of the gene, and no other mutations were found in the rest of the gene. The failure to recover a PCR product led to the conclusion that *bop5-3* has a

¹Using the new nomenclature for dynein genes in *Chlamydomonas* (Hom *et al.*, submitted), the IC138 subunit of the I1 inner arm dynein is encoded by the *DIC4* gene (previously known as the *BOP5* gene), and mutations in *DIC4* are designated *bop5* alleles.

Strain (CC #)	Motility	Axonemal phenotype	Original study
Control strains			
Wild-type <i>mt</i> + (CC-125)	Fast forward	Wild type	
Wild-type <i>mt</i> - (CC-124)	Fast forward	Wild type	
ppr2	Slow forward, no photoshock response	Defective CAV2 channel	Fujiu <i>et al.</i> , 2009
mia1 (CC-4265)	Slow forward, photo- taxis defect	hyperphosphorylation of IC138, missing Mia1p	King and Dutcher, 1997
mia2 (CC-4266)	Slow forward, photo- taxis defect	Hyperphosphorylation of IC138, missing Mia2p	King and Dutcher, 1997
mbo2 (CC-3664)	Backward swimming	Missing beak proteins, MBO2	Segal <i>et al.</i> , 1984; Tam and Lefebvre, 2002
pf10 (CC-3666)	Spins in circles	Unknown	Dutcher et al., 1988
uni1-2 (CC-2507)	Spins in circles	Uni-flagellate	Huang <i>et al.</i> , 1982
	·	-	
			Muster et al. 1007
$p_{1} = 3 (CC - 39 3)$	Slow forward	Missing II dynein	Myster et al., 1997
pt9-3::DHC1-G4 Ta (CC-3920)	Faster forward	Missing Ta HC motor domain	Nyster et al., 1999
p128; p19-2::DHC1 G4 (CC-3917)	Slow forward	domain	Myster et al., 1999
pf28; pf9-2 E8 (CC-3903)	Spins in circles	Missing outer arms, I1 dynein, full-length flagella	Porter <i>et al.</i> , 1992
ida2-6 (CC-3922)	Slow forward	Missing I1 dynein	Perrone et al., 2000
ida2-7 (CC-3923)	Slow forward	Missing I1 dynein	Perrone et al., 2000
ida2-7::DHC10 (CC-4401)	Faster forward	Missing 1 β HC motor domain	Perrone et al., 2000
pf28; ida2-7::DHC10 (CC-4079)	Slow forward	Missing outer arms, 1β motor domain	Perrone et al., 2000
IC138 mutants			
bop5-1 (CC-4080)	Slow forward	Truncated IC138, missing LC7b, FAP120	Dutcher et al., 1988; Hendrickson et al., 2004; Ikeda et al., 2009
bop5-2 mt+ (CC-4284)	Mixed motility	Missing IC138 subcomplex, MBO2	Bower et al., 2009
bop5-3 mt+ (CC-4389)	Slow forward	Missing IC138 subcomplex	Ikeda <i>et al.,</i> 2009
bop5-3 mt- (CC-4390)	Slow forward	Missing IC138 subcomplex	Ikeda <i>et al.,</i> 2009
bop5-4 mt-(CC-4391)	Slow forward	Missing IC138 subcomplex	lkeda <i>et al.</i> , 2009
bop5-5 mt+ (CC-4392)	Slow forward	Missing IC138 subcomplex	Ikeda <i>et al.,</i> 2009
bop5-6 mt+ (CC-4393)	Slow forward	Reduced IC138 subcomplex	lkeda et al., 2009
bop5-1::IC138 (CC-4081)	Fast forward	Wild type	Hendrickson <i>et al.</i> , 2004
bop5-2::IC138 mt+ (CC-4285)	Mostly backward	Missing MBO2	Bower et al., 2009
bop5-3::IC138 2D1 mt+ (CC-4395)	Fast forward	Wild type	This study
bop5-3::IC138 2E1 mt+	Fast forward	Wild type	This study
Double mutants			
pf10: bop5-3 (CC-4398)	Slow forward	Missing IC138 subcompley	This study
mbo2: $bop5-3$ (5e) (CC-4397)	Slow backward	Missing IC138 subcomplex MBO2n	This study
mbo2: bop5-3 (7b)	Slow backward	Missing IC138 subcomplex_MBO2p	This study
uni1-2: bop5-3 (CC-4399)	Uniflagellate spin	Missing IC138 subcomplex	This study
mia1: bop5-1 (CC-4394)	Slow forward	Truncated IC138 missing I C7b FAP120	This study
		Mia1p	
mia2; bop5-3 (CC-4396)	Slow forward	Missing IC138 subcomplex, Mia2p	This study
mia1; bop5-4 (CC-4400)	Slow forward	Missing IC138 subcomplex, Mia1p	This study
uni1-2; ida3	Uniflagellate, spin	Missing I1 dynein	Bayly et al., 2010

 TABLE 1: Strains used in this study.



FIGURE 2: Schematic diagram of the *bop5* mutations. The molecular defects in each *bop5* mutant are shown in this intron–exon diagram of the *IC138* (*DIC4*) gene. The *bop5-1* and *bop5-2* mutations were characterized in Hendrickson *et al.* (2004) and Bower *et al.* (2009). *bop5-3, 4, 5, 6* mutations were identified by PCR and sequencing of the *IC138* gene. *Bop5-3* has a deletion affecting the 5' end of the gene; *bop5-4* and *bop5-5* have mutations that result in premature stops; and *bop5-6* has a mutation in an intron that results in alternative splicing.

deletion that removes most of the 5'UTR and the first four exons but does not extend into the adjacent gene. The deleted region is between 1500 and 2000 base pairs (Figure 2 and unpublished data). bop5-4 contains a G-to-T mutation toward the end of the fourth exon that results in a premature stop codon (TAG) instead of a glutamic acid residue (GAG) (Figure 2). Another premature stop is created in bop5-5, from deletion of a single G that results in a frame-shift mutation and a stop codon at the end of the second exon (Figure 2). The last bop5 allele, bop5-6, contains a single G-to-A mutation in the third intron (Figure 2). This mutation does not directly alter the predicted wild-type splice site, but sequence analysis of RT-PCR products indicates that this mutation results in multiple alternatively spliced transcripts (unpublished data). Most of these transcripts contain premature stop codons and would likely not produce a functional protein. A small amount of normally spliced message is also present in bop5-6, which results in the expression of an IC138 polypeptide at lower levels than in wild type (Ikeda et al., 2009; Figure 3).

The various bop5 mutations affect expression of *IC138* and assembly of several other 11 subunits into the axoneme. Figure 3 shows an immunoblot that summarizes these results. As previously described (Hendrickson *et al.*, 2004; Bower *et al.*, 2009, Ikeda *et al.*, 2009), bop5-1 assembles a truncated IC138 and no FAP120 into the axoneme. Assembly of IC97 is not significantly affected. Similar to bop5-2, the bop5-3, bop5-4, and bop5-5 strains lack IC138, and do not assemble IC97 or FAP120 into the axoneme. bop5-6 shows a reduction of IC138, IC97, and FAP120 expression in the axoneme. IC140 serves as a loading control, as it is an 11 component the assembly of which is not affected by bop5 mutations (Bower *et al.*, 2009; Ikeda *et al.*, 2009). bop5-3 transformed with a wild-type copy of the *IC138* gene shows restoration of all 11

subunits missing in the *bop5-3* mutant. Because *bop5* mutants still assemble the two I1 dynein HCs, IC140, and other LCs (Bower *et al.*, 2009; Ikeda *et al.*, 2009), these mutants provide a unique opportunity to study the role of IC/LC subcomplexes in the regulation of dynein motor activity.

Forward swim speeds and microtubule sliding velocities are defective in bop5 mutants

Many I1 mutants, such as pf9, ida7, and ida2, show defects in swimming (the cells swim slowly, but smoothly, in a forward direction). These defects have led to the hypothesis that slow, smooth swimming is a hallmark of strains with defects in the I1 dynein complex (Myster et al., 1997; Perrone et al., 1998; 2000). Consistent with this hypothesis, the bop5 strains, which are not complicated by second mutations, also display a slow, smooth swimming phenotype (Figure 4A). The pf9-3 mutant swims most slowly (48.3 \pm 10.8 μ m/s), whereas bop5-1, bop5-3, bop5-4, and bop5-6 all swim between 70 and 90 μ m/s (73.8 \pm 12.7, 90.2 \pm 14.9, 77.1 \pm 11.2, and 78.5 \pm 11.2 µm/s, respectively). Neither bop5-2 nor bop5-5 was analyzed in greater detail as they both contain additional mutations that affect motility (Bower et al., 2009 and un-

published results). Transformation of *bop5-3* with the wild-type *IC138* gene restores forward swim speeds to near wild-type velocities (see Figure 4A).

We also investigated the phenotype of a *bop5-3*; *pf10* double mutant, as the original *bop5-1* mutant, which encodes a truncated IC138 polypeptide, was initially isolated as a suppressor of *pf10* (Dutcher *et al.*, 1988). Interestingly, the *bop5-3*; *pf10* double mutant, which lacks the IC138 subcomplex, also suppresses the *pf10* phenotype, as seen by tracking the forward motility of several cells in long exposure movies. Motility in the double mutants is not restored to wild-type levels, but the double mutant cells progress forward more rapidly than do *pf10* cells, which circle in place (Figure 4B).



FIGURE 3: Assembly of 11 dynein subunits in *bop5* mutant axonemes. Western blots of isolated axonemes were probed with antibodies to 11 dynein subunits. IC140 serves as a loading control, as it is not affected in any *bop5* mutant. IC138 is shifted in *bop5-1* and missing in *bop5-2* through *bop5-5*. FAP120 is missing in all *bop5* strains. IC138, IC97, and FAP120 are reduced in *bop5-6*. Note the reassembly of all 11 subunits in *bop5-3* strains that had been transformed with a wild-type copy of the *IC138* gene (2D1 and 2E1).



FIGURE 4: Forward swimming behavior and microtubule sliding in bop5 mutants. (A) Graph showing the average forward swimming velocities for wild type, pf9-3, bop5-1, bop5-3, bop5-4, bop5-6, and the bop5-3 rescued strains. At least 190–552 cells were measured for each strain. The swimming velocities of the IC138 rescued strains are significantly faster than the *bop5* mutants (p < 0.005), not significantly different from one another (p = 0.832), and only slightly slower than wild type (p < 0.005). The small differences (~5%) observed between wild type and rescued strains are typical of those seen with other transgene rescues. (B) Long exposure (1 s) images showing the forward progression of wild type, bop5-3, pf10, and bop5-3 pf10 cells. (C) Isolated axonemes from wild type and mutant strains were treated with protease and ATP, and the velocities of microtubule sliding were measuring during axoneme disintegration in vitro (n > 16 axonemes per strain). The sliding velocities of the bop5 strains are not significantly different from another or ida2-6, with the exception that bop5-4 is slightly faster than bop5-1 (p < 0.005). The sliding velocities of the IC138 rescued strains are significantly faster than all of the bop5 mutants, including bop5-3 (p < 0.005), but not significantly different from one another (p = 0.678) or wild type (p = 0.007 for 2D1 and p = 0.036 for 2E1).

Previous studies of 11 function showed that mutations in 11 subunits cause microtubule sliding defects (DiBella *et al.*, 2004b, Hendrickson *et al.*, 2004; Bower *et al.*, 2009; Wirschell *et al.*, 2009;

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Toba et al., 2011). Specifically, both bop5-1 and bop5-2 have reduced microtubule sliding velocities (Hendrickson et al., 2004; Bower et al., 2009). As expected, bop5-3, bop5-4, and bop5-6 also show decreases in microtubule sliding velocity, to ~75% of the wildtype rate and similar to that of the I1 mutant *ida2-6* (Figure 4C). Microtubule sliding velocities are restored to near wild-type rates in the bop5-3 IC138 rescued strains (Figure 4C).

Analysis of bop5-3 waveforms identifies defects in shear amplitude, sliding velocities, and bend propagation

Because bop5-1 results in the assembly of a truncated IC138, and bop5-2 has other mutations that affect motility (Hendrickson et al., 2004; Bower et al., 2009), the in vivo motility phenotypes of an IC138 null mutant have never been analyzed in significant detail. To investigate the IC138 null phenotypes, we first looked at the waveforms of forward swimming cells. Because bop5 mutants swim slowly but do not show a significant reduction in beat frequency (Table 2 and data not shown), and the ida3 mutant, which lacks the entire I1 complex, has an altered waveform (Bayly et al., 2010), we hypothesized that bop5-3 would also show changes in waveform. To better illustrate the parameters assayed, two individual waveform tracings from wild type and *ida3* are highlighted (Figure 5A). The principal bend is the bend with higher curvature, meaning the radius of a circle approximating the curve is smaller, and the reverse bend is the one with lower curvature. Although the waveform tracings from wild type and ida3 have similar maximum and minimum curvatures, the two waveforms look different because the regions of high curvature are of different sizes. If the curvature were imposed by two circles, as shown in Figure 5A, the ida3 flagellum would be wrapped around a shorter arc of each circle than the wild-type flagellum would be (Figure 5A). In other words, the bends (the regions of roughly constant, higher curvature) occur over smaller sections in ida3 flagella compared with wild type (Figure 5A). When the bends are shorter, the result is a narrower ("stunted") waveform, as in ida3 compared with wild type. Reducing the length of the bends reduces shear amplitude and shortens the stroke. This idea is also illustrated in the plots of the shear angle of each waveform, in which the regions of constant slope correspond to portions of the flagella with constant curvature. In the shear angle plots of wild-type flagella, relatively long sections have either a constant negative or constant positive slope. These regions of constant slope are shorter in shear angle plots for ida3 flagella, so the curves have lower amplitude, and the distribution of shear angles is narrower in ida3 flagella (Figure 5A).

The differences between bop5-3, ida3, and wild type are evident in the traces of the overall waveform beat pattern (Figure 5B): bop5-3 waveforms are narrower than wild type, but not as severely affected as ida3. These differences can also be seen in the supplemental movies (Supplemental Movies S1-S3). To characterize the waveforms quantitatively, shear angle curves were plotted (Figure 5C), and various parameters of the curves were calculated, as described in Bayly et al., 2010. Qualitatively, wild type exhibits larger oscillations in shear angle than does bop5-3, and the bop5-3 shear angle curves have larger oscillations than those of ida3 (Figure 5C); the bop5-3 and ida3 graphs of shear angle distribution are narrower than wild type. These oscillations in shear angle are represented quantitatively by the root mean square shear amplitude. Wild-type values are higher than those of bop5-3, and bop5-3 values are higher than those of ida3 (0.68 \pm 0.05, 0.58 ± 0.05 , and 0.52 ± 0.04 rad, respectively, p < 0.05; Table 2).

The sliding velocities in the principal and reverse bends (average minimum, or R, sliding velocity and average maximum, or P, sliding velocity) in *bop5-3* and *ida3* also differ significantly from their values

	Wild type	bop5-3	ida3
Flagellar beat frequency (Hz)	69.45 ± 8.41	69.59 ± 10.35	64.29 ± 7.26
Body revolutions per second*†	4.52 ± 0.95	3.55 ± 0.91	2.24 ± 0.40
Length (µm)*†	12.79 ± 1.15	10.34 ± 2.14	13.87 ± 1.17
Avg. root mean square shear amplitude (rad)* †	0.68 ± 0.05	0.58 ± 0.05	0.52 ± 0.04
Avg. min. (R) sliding velocity-physical (rad/s)*	-404.1 ± 50.06	-340.2 ± 51.09	-313.3 ± 35.73
Avg. max. (P) sliding velocity-physical (rad/s)*†	543.6 ± 54.77	488.7 ± 38.23	430.8 ± 42.04
Ratio of propagation speeds (Reverse/Principal)* †	1.03 ± 0.10	0.87 ± 0.16	0.76 ± 0.10
Avg. distance between principal and reverse curva-	0.75 ± 0.09	0.78 ± 0.04	0.72 ± 0.08

*Parameters for which values for bop5-3 and ida3 were significantly different from wild type.

[†]Parameters for which values for *bop5-3* and *ida3* were significantly different from each other (p < 0.05). The average minimum and maximum sliding velocities are spatial averages of the minimum (maximum) sliding velocity at each point on the flagellum, averaged over the middle 80% of its length. The minimum sliding velocity occurs in the reverse (R) bend and characterizes the effective stroke; the maximum sliding velocity occurs in the principal (P) bend and describes the recovery stroke.

TABLE 2: Flagellar waveform parameters.

in wild type. These parameters describe the average minimum (or maximum) interdoublet sliding velocity at each point over the middle 80% of the flagellar length. The minimal sliding velocity at a given point occurs as the reverse bend propagates past that point; the maximum sliding velocity at a point occurs as the principal bend propagates. Both the *bop5-3* and *ida3* display reduced sliding velocities in the reverse bend relative to wild type, but the values for *bop5-3* and *ida3* are not significantly different from each other. The sliding velocity of the principal bend in *bop5-3* is significantly lower than in wild type but significantly higher than in *ida3* (Table 2). These reductions in the calculated microtubule sliding velocities are similar to the reductions measured in the in vitro sliding assays (Figure 4C).

The final parameter that exhibits differences between *bop5-3* and wild type is the propagation speed of the reverse bend (Table 2). As this value is also dependent on flagellar length, we compensated for minor variations in length by analyzing the ratio of the reverse propagation speed to the principal propagation speed. The values of this ratio in *bop5-3* and *ida3* are 15–20% lower than those of wild type. These lower values result primarily from the lower propagation speed of the reverse bend, suggesting that the effective stroke is affected most strongly. Speed of bend propagation is closely related to (although not completely determined by) the speed of microtubule sliding, which is significantly lower during the effective strokes of *bop5-3* and *ida3* (see preceding paragraph and *Discussion*).

I1 dynein mutants exhibit defects in photoshock behavior

To further characterize the effect of the missing IC138 subcomplex, we compared the photoshock response in the 11 mutant strains to that of wild type and other mutants (Table 3). Wild-type cells normally swim forward using an asymmetric ciliary-type waveform, but abruptly halt upon sudden exposure to bright light and convert their waveforms to a more symmetric flagellar-type beat in response to elevated intracellular calcium. This calcium increase results in a brief period (~0.5 s) of backward movement (photoshock), but then, as intracellular calcium levels drop, the cells switch back to the more asymmetric ciliary waveform and resume normal forward motility (Witman, 1993; see Table 3 and Figure 6). In contrast, mutant cells with defects in the flagellar membrane voltage activated calcium channel (*ppr2*) or mutant cells with defects in outer doublet beak structures (*mbo2*) do not alter their motility in response to sudden

bright light (Segal *et al.*, 1984; Matsuda *et al.*, 1998; Fujiu *et al.*, 2009; see Table 3). All of the 11 mutants, however, displayed an altered photoshock response, where the cells paused upon bright light exposure (suggesting that they detected the light stimulus), but the cells did not appear to swim backward any significant distance (Figure 6 and Table 3). The *mia1* and *mia2* mutants, which exhibit altered IC138 phosphorylation (King and Dutcher, 1997) also did not appear to reverse during the photoshock response. Interestingly, backward movement during photoshock can be observed in *bop5-3*, *bop5-1*, *pf9-3*, and *ida2-7* strains that have been rescued by transformation with genes encoding IC138, 1 α HC, or 1 β HC, respectively (Table 3).

One possible interpretation is that cells with an altered 11 complex might not be capable of converting their waveforms to the more symmetric flagellar-type beat. Previous analysis of an 11 mutant has shown, however, that waveform conversions are possible (Brokaw and Kamiya, 1987). In addition, we constructed double mutants containing both *bop5-3* and *mbo2* mutations, and we observed vigorous backward swimming when the double mutant cells were viewed with conventional light microscopy (Table 1). These results strongly suggest that cells lacking the IC138 subcomplex can alter their waveforms, but may not generate enough power to swim backward efficiently during photoshock (see *Discussion*).

I1 null mutants, but not bop5 mutants, have impaired phototaxis

Because I1 mutants such as *pf9/ida1*, *ida2*, *ida3*, *mia1*, and *mia2* exhibit phototaxis defects (King and Dutcher, 1997; Okita *et al.*, 2005), we assayed phototactic ability in the *bop5* mutants using both photoaccumulation and quantitative phototaxis assays (Table 4, Figure 7, and Supplemental Movie S4). As the phototaxis measurements showed similar results to the photoaccumulation assays (Table 4), phototaxis diagrams are illustrated for only a subset of the strains (Figure 7B). Wild-type mating-type plus (CC-125) and mating-type minus (CC-124) cells showed positive and negative taxis, as reported previously (Pazour *et al.*, 1995; Okita *et al.*, 2005). The phototactic sign (positive or negative) depends on the genetic background (*AGG1* or *agg1*) of each strain (Smyth and Ebersold, 1985; Iomini *et al.*, 2006). As expected, the I1 DHC mutants (*pf9-3* and *ida2-7*) did not show significant phototaxis in either direction (Myster *et al.*, 1999; Perrone *et al.*, 2000). The DHC-rescued strains



were able to phototax, although the strain with the full-length 1 β DHC (*pf9-3::DHC1*) displayed more robust photoaccumulation than the strain with the full-length 1 α DHC (*ida2-7::DHC10*) (Table 4).

(pf9-3::DHC1 and ida2-7::DHC10), however,

The mia1 and mia2 mutants, which contain the entire I1 complex but exhibit hyperphosphorylation of IC138, are not able to phototax (King and Dutcher 1997; Okita et al., 2005). Surprisingly, all of the bop5 mutant strains were able to phototax, in either the positive or negative direction, nearly as well as wild type (Figure 7 and Table 4). Because previous studies have suggested that hyperphosphorylation of IC138 might be the basis of the phototaxis defects in the mia mutants, we wondered if loss of the IC138 subcomplex could suppress the mia phenotype. To test this hypothesis, we constructed double mutant strains containing bop5-1, bop5-3, or bop5-4 with either mia1 or mia2. Interestingly, the bop5; mia double mutants were not able to phototax, and their forward swimming velocities were similar to those displayed by the single mia mutants (Table 4, Figure 7, and unpublished data).

DISCUSSION

The IC138 subcomplex is necessary for normal flagellar motility

Although the IC138 subcomplex is not required for the stability, targeting, and/or docking of I1 dynein in the axoneme (Bower et al., 2009), the IC138 subcomplex is essential for control of the size and shape of the flagellar waveform during forward swimming. All four bop5 mutant strains (bop5-1, bop5-3, bop5-4, bop5-6) show similar motility defects, suggesting not only that the IC138 subcomplex is necessary for normal flagellar motility, but that either a reduction in IC138 subcomplex subunits (bop5-6) or a truncated IC138 and missing LC7b (bop5-1) can also cause abnormal motility (Hendrickson et al., 2004; Bower et al., 2009). All four mutants swim at ~2/3 the velocity of wildtype strains (Figure 4A). The reduction in swimming velocity is caused by altered and inefficient flagellar waveforms, not reduced beat frequency, similar to other I1 mutants (Brokaw and Kamiya, 1987; Perrone et al.,

ida3, showing every $1/12^{\text{th}}$ cycle. (C) Spatial distribution of shear angle of the same five representative cells of wild type, *bop5-3*, and *ida3*, showing every $1/12^{\text{th}}$ of a cycle. Data on wild type and *ida3* are reprinted with modification from Bayly et *al.* (2010).

FIGURE 5: Asymmetric waveforms of 11 mutant strains during forward swimming. (A) Illustration of flagellar waveforms (left) and shear angle curves (right) plotted vs. normalized distance along the flagellum (*s/L*). The effects of bend curvature and bend arc length on the amplitude of flagellar waveform are illustrated on the left. Either reduced curvature (larger radius of curvature) or shorter curved regions (less wrapping) will lead to narrower waveforms. The effective stroke is dominated by the reverse bend, and the recovery stroke is dominated by the principal bend. (B) Flagellar waveforms of five representative cells of wild type, *bop5-3*, and

Strain	Reversal (%)	Pause/direction change (%)	No change (%)	Number of cells scored
Wild-type +	92	8	0	50
Wild-type –	90	10	0	30
mia1-1	0	100	0	30
mia2-1	0	100	0	30
mbo2	0	5	95	20
ppr2	0	12	88	25
pf9-3	0	79	21	53
pf9-3::DHC1	90	10	0	30
pf28; pf9-2::DHC1	0	45	55	78
pf28; pf9-2 E8	0	52	48	66
ida2-7	0	81	19	67
ida2-7::DHC10	16	66	18	85
pf28; ida2-7::DHC10	0	57	43	23
bop5-1	0	80	20	50
bop5-1::IC138	70	27	3	30
bop5-3 mt+	0	100	0	54
bop5-3::IC138 (2D1)	63	37	0	30
bop5-3::IC138 (2E1)	67	33	0	30
bop5-3; mbo2 (5e)	0	0	100	30
bop5-3; mbo2 (7b)	0	0	100	20
bop5-4	0	97	3	30
bop5-5	0	68	32	30
bop5-6	0	97	3	30

Actively moving cells were exposed to a brief flash of bright light and then scored for reversal, pausing, and/or change in forward direction, or no change in direction of movement. The most prevalent phenotypes are highlighted in bold type, with the exception of ida2-7::DHC10 (see text for details).

TABLE 3: Photoshock responses in wild type and mutant strains.

2000; Myster et al., 1997, 1999; Bayly et al., 2010). The reduced swim speeds are further correlated with reduced microtubule sliding velocities in *bop5* axonemes (Figure 4C). In addition, the *bop5* mutations partially suppress the motility defects seen in the *pf10* mutant (Figure 4B). These observations suggest that the IC138 subcomplex and the protein product encoded by the *PF10* gene interact directly or indirectly to regulate motility. Further insight into this interaction will require identification and localization of the *PF10* gene product.

Loss of the IC138 subcomplex changes the shape and amplitude of the flagellar waveform

Comparisons of waveform parameters between wild-type, *bop5-3*, and *ida3* mutants have provided new insights into the role of both the 11 dynein and the IC138 subcomplex in controlling motility. First, the overall distances between principle and reverse curvature extremes are approximately equal in all three strains (Table 2 and Figure 5B). Strains lacking the IC138 subcomplex or the entire 11 dynein are still able to generate bends with curvatures similar to wild type, suggesting that the net forces bending the flagella are similar to those in wild type (i.e., that the loss of the I1 dynein or IC138 subcomplex does not dramatically affect the total dynein force summed over the length of the flagellum).

In contrast, the calculated microtubule sliding velocities in mutants lacking I1 or the IC138 subcomplex significantly differ from

the sliding velocities in wild-type flagella (Table 2). Moreover, calculated sliding velocity and shear amplitude are affected more severely in *ida3* than in *bop5-3* flagella, suggesting that the 11 dynein contributes to control of microtubule sliding and shear, even in the absence of the IC138 subcomplex (Table 2). Consistent with the calculated reduction in waveform sliding velocity, direct measurement in isolated axonemes also reveals reductions in the microtubule sliding velocity when the IC138 subcomplex fails to assemble (Figure 4C). This reduction in sliding velocity suggests that, even if the total force summed along the length of the flagellum is similar to the corresponding cumulative force in wild type, the local force generated by dyneins at any one point on the flagellum is reduced in I1 mutants.

If dynein force generation is reduced at any one point, but similar overall, we wondered how dynein force generation is being altered in the 11 mutants. The spatial distribution of dynein activity must be different, and this difference affects the overall shape of the mutant waveforms, as seen by the reduction in shear amplitude (Table 2). It is clear that *ida3* and *bop5-3* flagella exhibit shorter regions of constant curvature than do wild type. Once again, the effect is most severe in *ida3*, and intermediate in *bop5-3*. This may mean that fewer dynein cross-bridges are active per unit length of the flagellum than in wild type or, alternatively, that the mechanisms that coordinate the selective activation and inactivation of different dyneins during bending are disrupted in 11 mutants. The latter may



FIGURE 6: Photoshock behavior in 11 mutant strains. The movement of wild type, *ppr2*, and *bop5-3* cells were recorded before, during, and after exposure to a bright light stimulus. Six frames from each movie are shown here, illustrating the movement of an individual cell for ~0.90 s. Forward motion is toward the top of the figure, and reverse motion is toward the bottom. On sudden exposure to bright light, the wild-type cell stops forward motion, moves backward briefly, and then resumes forward motion. The *ppr2* cell moves forward at a constant rate throughout the assay. The *bop5-3* cell stops briefly, but does not reverse before recovering and swimming forward again.

be consistent with the notion that I1 dynein plays a role in the local inhibition of sliding driven by other dyneins (Porter *et al.*, 1992; Smith, 2002; Kotani *et al.*, 2007; Toba *et al.*, 2011). Moreover, multiple connections between I1 dynein and both outer and inner arm dyneins have been observed by cryoelectron tomography of isolated axonemes (Nicastro *et al.*, 2006; Bui *et al.*, 2008; Heuser *et al.*, 2009).

Another parameter that differs significantly between wild type and the I1 mutants is the ratio of propagation speeds. This ratio represents the speed of the reverse bend compared with the speed of the principal bend and reveals if the dyneins in one type of bend are activated/inactivated differently than in another type of bend. The ratio of propagation speed is reduced more severely in ida3 flagella than in bop5-3 flagella, but both ratios are lower than the ratio for wild-type flagella. The reduction is caused by the reduced speed of the reverse bend (see Table 2). One possible explanation is that dyneins are not being properly inactivated during the effective stroke. The lack of inactivation may limit curve progression, which could slow down the overall speed of the bend. The reduction of both shear amplitude and reverse bend propagation speed in bop5 mutants leads to the hypothesis that dyneins are not being properly activated and inactivated temporally and spatially along the flagella, which is consistent with the model that the IC138 subcomplex is important in regulating shape of the flagellar bend (Hendrickson et al., 2004; Bower et al., 2009). The defects in waveform also explain the slow swimming phenotype seen in both bop5 and other I1 mutants.

Strain	Photoaccumulation	Phototaxis	Direction
Wild-type <i>mt</i> +	Yes	Yes	Positive
Wild-type <i>mt</i> -	Yes	Yes	Negative
mia1	No	No	NA
mia2	No	No	NA
ppr2	Yes	NT	Positive
pf9-3	No	NT	NA
pf9-3::DHC1 G4 1a	Yes	NT	Negative
pf28; pf9-2::DHC1 G4	No	NT	NA
pf28; pf9-2 E8	No	NT	NA
ida2-7	No	NT	NA
ida2-7::DHC10	Yes, weak	NT	Positive
pf28; ida2-7::DHC10	No	NT	NA
bop5-1	Yes	Yes	Positive
bop5-1::IC138	Yes	NT	Positive
bop5-3 mt+	Yes	Yes	Negative
bop5-3::IC138 (2D1)	Yes	Yes	Negative
bop5-3::IC138 (2E1)	Yes	Yes	Negative
bop5-4	Yes	Yes	Positive
bop5-6	Yes	Yes	Variable
mia1; bop5-1	No	NT	NA
mia2; bop5-3	No	No	NA
mia1; bop5-4	No	No	NA

NT, not tested in this study; NA, not applicable.

TABLE 4: Phototaxis responses in wild-type and mutant strains.



FIGURE 7: Phototaxis behavior in wild type and I1 mutant strains. (A) Photographs of selected strains showing their photoaccumulation behavior in a Petri dish after 10 min in front of a light box. The light box is located at the right of the page. (B) Polar histograms showing the swimming direction of individual cells ~5 s after the phototactic light source was turned on at 0 degrees (right side).

The IC138 subcomplex is necessary for an efficient photophobic response

One important feature of *Chlamydomonas* behavior is the photoshock response. All 11 mutants showed a similar defect in the photoshock response: They paused, but did not reverse (Figure 6 and Table 3). This includes the *mia* mutants, which have altered phosphorylated IC138 levels, and *bop5-1*, which is only missing part of IC138, LC7b, and FAP120 (Table 3). The defects in the photoshock response are partially restored in the rescued strains. Interestingly, in mutant cells missing the 1 α motor domain, the photoshock response is more effectively restored compared with mutants lacking only the 1 β motor domain (Table 3). These results suggest that the 1 β motor domain is more important in regulating the photoshock response. This hypothesis is in line with recent work that suggests that the 1 β HC is a more effective motor that the 1 α HC (Toba *et al.*, 2011).

The basis of the "no reversal" phenotype is not well understood, but it is consistent with the phenotype seen in other dynein mutants (Kamiya and Okamoto, 1985; King and Kamiya, 2009). One hypothesis is that the cells may not be able to generate a backward, more symmetric, flagellar-type waveform without assembly of 11 dynein. *bop5-3; mbo2* mutant cells, however, swim backward efficiently (Table 3), and waveform conversion has been detected in the 11 mutant, *ida98*, now known as *ida1-1* (Brokaw and Kamiya, 1987), so this seems unlikely. It is more probable that, although *bop5* cells can switch their waveforms, these waveforms do not generate enough power to result in the net backward movement of the cell body during the brief time period (~0.5 s) of the photoshock response.

The IC138 subcomplex is not required for phototaxis

Phototaxis is a sensitive measure of the coordination and precise control of bending for each flagellum in *Chlamydomonas* (Witman, 1993). I1 mutant cells display defects in phototaxis when measured using simple photoaccumulation assays (King and Dutcher, 1997) or more quantitative phototaxis measurements (Okita *et al.*, 2005). We have confirmed these observations using the I1 DHC mutants (Table 4; Myster *et al.*, 1999; Perrone *et al.*, 2000). Moreover, assembly of only one of the two motor head domains is sufficient to restore phototaxis ability, as the loss of either the 1α or 1β motor domain does not impair phototaxis (Table 4; Myster *et al.*, 1999;

Perrone et al., 2000). Cells missing the 1α motor domain show better phototaxis than do cells lacking the 1β motor domain (see Table 4). These results correlate with recent observations showing that the 1β dynein has higher motor activity than the 1α dynein (Toba et al., 2011). Although the 11 motor domain mutants and outer arm mutants (*pf28*) can each undergo phototaxis, double mutant cells missing both the outer arms and an 11 motor domain cannot (Table 4). This observation suggests a functional interaction between the motor domains in 11 dynein and the outer dynein arms.

Because nearly all of the *bop5* mutant strains can phototax, we conclude that the IC138 subcomplex is not absolutely required for phototaxis (Table 4). These observations suggest that the remaining subunits of the I1 complex must play an essential role in phototaxis, even in the absence of the IC138 subcomplex. The *bop5; mia* double mutant

strains, however, are all defective in phototaxis (Table 4 and Figure 7). This observation is significant because it has been assumed that phototaxis defects in the *mia* mutants were primarily the result of hyperphosphorylation of IC138, leading to inhibition of I1 dynein. Because the rest of the I1 dynein is present, but the IC138 subcomplex is missing in the double mutants, we postulate that defects in other, unknown axonemal proteins may be the real cause of the phototaxis defects observed in the *mia* mutants.

The C-terminal region of IC138 plays a critical role in I1 dynein function

The motility defects of the *bop5-1* (truncated IC138) mutant are similar to those of the *bop5* null strains. The forward swimming velocities of *bop5-1*, *bop5-3*, *bop5-4*, and *bop5-6* are ~80 µm/s, which suggests that their waveform parameters may be similar (Figure 4A). The *bop5-1* strain also displays a similar defect in the photoshock response (Table 3). These results indicate that the C-terminal region of IC138, which binds LC7b and possibly FAP120, plays a critical role in I1 dynein function (Hendrickson *et al.*, 2004; Ikeda *et al.*, 2009).

Our study of the bop5 mutant alleles provides new insights into the function of the IC138 subcomplex. Loss of the IC138 subcomplex has several effects in vivo: reduced swimming velocities, altered waveforms, and inefficient reversal responses during photoshock. Microtubule sliding velocities are reduced both in vivo and during sliding disintegration in vitro. The net result is that the amplitude of the flagellar waveform is reduced: bop5 cells are less efficient at moving forward during normal motility and less effective at moving backward during photoshock. Contrary to expectation, however, loss of the IC138 subcomplex does not result in phototaxis defects. More work is needed to determine exactly how the I1 dynein contributes to control of phototaxis without the IC138 subcomplex. Nonetheless, evidence presented here demonstrates that the IC138 subcomplex plays an important role in controlling both the shape of the flagellar waveform and the response of the cell to environmental stimuli. Given the high degree of conservation of I1 dynein subunits (Wickstead and Gull, 2007), the IC138 subcomplex likely plays a critical role in regulating flagellar motility in many species. In addition, the bop5 mutants demonstrate that an IC/LC complex can modulate dynein motor activity independent of any effects on complex stability and/or cargo binding.

MATERIALS AND METHODS

Strains, culture conditions, and genetic analyses

All strains discussed in this study are shown in Table 1. Strains were grown on Tris acetate phosphate (TAP) medium (Harris, 1989). For most assays, a pea-sized amount of cells was resuspended in minimal medium lacking nitrogen (Harris, 1989) and rocked overnight under constant light to encourage maximum flagellar assembly and motility. Photobehaviors, especially phototaxis, were more robust in gametic versus vegetative cells. For flagellar waveform analysis, cells were cultured overnight in R medium (Harris, 1989).

Rescued *bop5-3* strains were generated by cotransformation of *bop5-3 mt+* with the pSI103 plasmid containing the *aphVIII* gene (Sizova *et al.*, 2001) and the pHX plasmid containing the gene that encodes IC138 (Hendrickson *et al.*, 2004). Transformants that grew on TAP plus 10 μ g/ml paromomycin were screened for rescue of the motility and protein defects. To generate double mutants, nonparental ditype tetrads were dissected and analyzed by light microscopy, PCR, and/or Western blots to identify the appropriate double mutant combination.

PCR analyses

PCR and sequencing of the *IC138* gene in each mutant were performed using primers designed in the MacVector program (MacVector, Cary, NC) to the *IC138* genomic sequence (GenBank accession AY743342). The PCR products were sequenced directly (Genewiz, South Plainfield, NJ), and specific mutations or deletions were identified in *bop5-3*, *bop5-4*, *bop5-5*, and *bop5-6*.

Isolation of axonemes, SDS–PAGE, and Western blot analysis

To isolate axonemes, cells were subjected to pH shock, and then isolated flagella were demembranated by 0.1% NP-40 treatment as described previously (Witman, 1986). The resulting axonemes were resuspended in HMEEN (10 mM HEPES, pH 7.4, 5 mM MgSO₄, 1 mM EGTA, 0.1 mM EDTA, 30 mM NaCl) with 0.1 μ g/ml protease inhibitors (pepstatin, leupeptin, aprotinin) and 1 mM dithiothreitol (Bower *et al.*, 2009).

Axoneme samples were run on 5–15% polyacrylamide gradient gels and transferred to Immobilon P (Millipore, Billerica, MA) as described previously (Bower *et al.*, 2009). Antibodies were used at the following dilutions (vol/vol): IC138, 1:10,000 (Hendrickson *et al.*, 2004); IC97, 1:10,000 (Wirschell *et al.*, 2009); IC140, 1:10,000 (Yang and Sale, 1998); FAP120, 1:10,000 (Ikeda *et al.*, 2009); IC69, 1:50,000 (Sigma-Aldrich, St. Louis, MO). Signal was detected with alkaline-phosphatase–conjugated secondary antibodies and the Tropix detection system (Bower *et al.*, 2009).

Analysis of swimming speed and motility assays

To measure forward swimming velocity, gametic cells were viewed on an Axioscope (Carl Zeiss, Thornwood, NY) with 20x phasecontrast optics and a halogen light source. Short (30 s – 1 min) movies were recorded with a Rolera-MGi EM-CCD camera (Q-Imaging, Tucson, AZ), and velocities were calculated using MetaMorph software (version 7.6.5.0; Molecular Devices, Sunnyvale, CA), as described previously (Bower *et al.*, 2009). All data are expressed as mean \pm SD. *F* tests and *t* tests were used to determine the significance of differences between means (Supplemental Table S1). To assay motility in a population of cells, long (1 s) exposures were taken with the Rolera-MGi EM-CCD camera. Final images are cropped regions of the field of view that show several motile cells.

Microtubule sliding assays

Microtubule sliding velocity was determined using the methods developed by Okagaki and Kamiya (1986) with minor modifications (Howard et al., 1994; Habermacher and Sale, 1996, 1997; Hendrickson et al., 2004; Toba et al., 2011). Flagella were resuspended in buffer lacking protease inhibitors, demembranated with 0.5% NP-40, and placed in perfusion chambers. Sliding was started by adding buffer with 1 mM ATP and 3 µg/ml subtilisin A type VIII protease (Sigma-Aldrich). Recordings of sliding were made on an Axiovert 35 microscope (Carl Zeiss) with dark field optics and a Silicon intensified camera (VE-1000; Dage-MTI, Michigan City, IN). The analogue videos were digitized, and velocity was determined manually on the flat computer screen by measuring displacement of microtubules calibrated with a stage micrometer. All data are expressed as mean \pm SD, and F tests and t tests were used to determine the significance of differences between means (Supplemental Table S2).

Flagellar waveform analysis of forward swimming cells

Analysis of the flagellar waveform during forward swimming was done using bop5-3; uni1 double mutant cells grown overnight in minimal medium. The acquisition and analysis of flagella waveforms were conducted as described in Bayly et al. (2010). Briefly, 600-frame videos (350 frames per second, 2.88 ms shutter speed) were taken on a Zeiss Universal Microscope (Carl Zeiss) at 100× phase contrast with a Dragonfly Express IEEE-1394b Digital Camera System and FlyCapture software (Point Grey Research, Scottsdale, AZ) and saved as avi files. Videos were analyzed using custom Matlab programs (The Mathworks, Natick, MA), which include correction of rigid-body motion, identification of flagellar point clouds, sorting of the point clouds, fitting of curves to the data, and characterization and statistical analysis of the final waveforms (Bayly et al., 2010). To determine if a waveform parameter is dependent on flagellar length, SAS 9.2 software (SAS Institute, Cary, NC) was used to create a general linear model of the data with the strain as a class variable (see also Supplemental Table S3).

High-speed movies of biflagellate cells were taken on an Axioscope (Carl Zeiss) with 63× DIC optics and a halogen light source. Movies were recorded with a 1024 PCI high-speed camera (Motion Engineering Company, Indianapolis, IN) at 500 frames per second. To facilitate viewing the movies, ImageJ (National Institutes of Heatlh, Bethesda, MD) was used to compile every fourth frame into a QuickTime movie with a playback speed of 15–30 frames per second, depending on the strain. Movies of wild type, *bop5-3*, and *pf9-3* are Supplemental Movies S1, S2, and S3.

Photoaccumulation, phototaxis, and photoshock assays

For photoaccumulation assays, gametic cells at a concentration of 2.5×10^6 cells/ml were transferred to 35 mm Petri dishes, and dark adapted for 10 min. Cells were then exposed to a fluorescent light source located 30 cm away (2 W/m²) for 10 min. Cell densities on both sides of the dish were determined at various time points and recorded by digital photography to determine relative photoaccumulation ability (Supplemental Movie S4). Each strain was independently assayed at least three times or tested separately for phototaxis to confirm the results. For phototaxis, cells were dark adapted in red light before illumination by unidirectional blue-green light. The cell behavior was recorded, and the angle between the light direction and the swimming direction of each cell was measured and pooled into 30° bins, as described previously (Okita *et al.*, 2005). Where applicable, the direction of the movement relative to the light source is noted: positive if toward the light source, and negative if away from the light source.

To assay for the photoshock response, we noted whether forward swimming cells exhibited any change in behavior upon brief exposure to bright light. Cell movement was recorded using phasecontrast microscopy at 20× and a red filter. The cells were then briefly exposed to bright light using a green fluorescent protein excitation filter on the epi-illuminescence port ($\lambda = 450-500$ nm). The position of each cell (n > 30) before and after light exposure was traced and scored as no change, paused, changed in forward direction only, or reversed.

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