

**REPORT**

# Ciliary force-responsive striated fibers promote basal body connections and cortical interactions

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**Multi-ciliary arrays promote fluid flow and cellular motility using the polarized and coordinated beating of hundreds of motile cilia. *Tetrahymena* basal bodies (BBs) nucleate and position cilia, whereby BB-associated striated fibers (SFs) promote BB anchorage and orientation into ciliary rows. Mutants that shorten SFs cause disoriented BBs. In contrast to the cytaxis model, we show that disoriented BBs with short SFs can regain normal orientation if SF length is restored. In addition, SFs adopt unique lengths by their shrinkage and growth to establish and maintain BB connections and cortical interactions in a ciliary force-dependent mechanism. *Tetrahymena* SFs comprise at least eight uniquely localizing proteins belonging to the SF-assemblin family. Loss of different proteins that localize to the SF base disrupts either SF steady-state length or ciliary force-induced SF elongation. Thus, the dynamic regulation of SFs promotes BB connections and cortical interactions to organize ciliary arrays.**

## Introduction

Multi-ciliary arrays comprise hundreds of hydrodynamically coupled cilia that beat in a coordinated and polarized manner. Basal bodies (BBs) nucleate, position, and anchor cilia at the cell cortex. Beating cilia produce both hydrodynamic flow across the cell surface and mechanical forces that are transduced to the BB anchors (Dirksen, 1971; Vernon and Woolley, 2004). Because of the asymmetric nature of ciliary beating, several forces are imposed upon BBs. These include oscillatory, compressive, and rotational forces during the power and recovery strokes (Cheung and Jahn, 1976; Narematsu et al., 2015; Riedel-Kruse et al., 2007). Nevertheless, BBs maintain their position and polar orientation.

Cilia-generated mechanical forces are resisted through BB connections with neighboring BBs and with the cell cortex. These linkages are promoted by BB appendage structures that are classified into distal appendages, basal feet, and striated fibers (SFs). BB distal appendages promote BB docking to the plasma membrane (Garcia and Reiter, 2016; Vertii et al., 2016). In amphibians, basal feet and SFs are polarized along the ciliary beat axis, but are oriented in opposite directions (Hard and Rieder, 1983; Werner et al., 2011). Both structures are generally thought to maintain BB position and orientation by mediating interactions with cortical microtubule, actin, and intermediate filament cytoskeletons (Antoniades et al., 2014; Kunimoto et al., 2012; Lemullois et al., 1987; Vladar et al., 2012). BB-associated SFs are striated structures that are conserved

across ciliated eukaryotes (Holberton et al., 1988; Lechtreck and Melkonian, 1991; Yang et al., 2002). In vertebrates, SFs consist of proteins proximal to the BB (C-Nap1, Centlein, and Cep68) that link to proteins that form the SF (Rootletin, Cep68, and Lrrc45; Fang et al., 2014; He et al., 2013; Vlijm et al., 2018). How BB-associated SFs respond to mechanical forces and interact with the cytoskeleton in multi-ciliated arrays remains poorly understood.

The BBs within *Tetrahymena* multi-ciliary arrays are arranged in longitudinal ciliary rows and possess microtubule appendages and SFs. The microtubule appendages consist of both the post-ciliary microtubule (pcMT) bundles, which project posteriorly from BBs, and the transverse microtubule bundles, which extend rightward (when viewed from the outside of the cell) to the adjacent ciliary row (Allen, 1967; Junker et al., 2019). SFs project anteriorly, to connect with anterior BBs by crossing the surface of the pcMT bundles (Allen, 1967; Junker et al., 2019; Pitelka, 1961). Ciliate cortical organization is promoted by both global and local polarity cues (Frankel, 1989; Frankel, 2008; Sonneborn, 1964). Cytotaxis is a local and nongenetic polarity mechanism, whereby preexisting BBs and their associated structures transmit local polarity information to guide the organization and orientation of new BBs (Beisson and Sonneborn, 1965; Frankel, 1964; Ng and Frankel, 1977; Sonneborn, 1964; Tartar, 1956). However, it is unclear whether and how the cell's

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cortical architecture recovers when the local polarity of the system is compromised.

The *Tetrahymena* BB orientation-defective mutant *disA-1* revealed that normal SF length is required for proper BB orientation (Galati et al., 2014; Jerka-Dziadosz et al., 1995). This facilitates BB connections within BB rows and enables the propagation of metachronal ciliary beating for cellular motility (Narematsu et al., 2015; Tamm, 1984; Tamm, 1999). However, the mechanisms by which SFs promote BB organization and orientation, and how their lengths are controlled, remain unknown.

Here, we show that *Tetrahymena thermophila* SFs physically link neighboring BBs to each other and to the cell cortex to organize, orient, and reorient BBs. SF lengths respond to changes in ciliary forces, such that elevated or reduced cilia-dependent forces cause SFs to elongate and shorten, respectively. As with vertebrate SFs, *T. thermophila* SFs are composed of a complex network of components that localize to different domains of the SF structure. Components localizing to the SF base ensure both (1) steady-state SF length and (2) elevated ciliary force-induced SF elongation. Using mutants of SF base components to separate these functions, we illuminate the important roles that the unique length states of SFs play in organizing and orienting BBs. These findings serve as a foundation for understanding the role of SF dynamics in anchoring BBs and hydrodynamic flow.

## Results and discussion

### SFs promote BB reorientation

Cells with severely disoriented BBs in *disA-1* mutants are rescued by the reintroduction of WT *DISA-1* (*DISA-1* rescue; Galati et al., 2014). Cytotaxis is a local and nongenetic polarity mechanism, whereby preexisting BBs guide the position and orientation of new BBs such that the existing cortical architecture is propagated to future generations (Beisson and Sonneborn, 1965; Ng and Frankel, 1977; Sonneborn, 1964; Tartar, 1956). However, in contrast to the cytaxis model, rescue of disoriented BBs in *DISA-1* rescue cells suggests that BBs retain the ability to correct their orientation, even in a landscape of disorganized BBs. To visualize BB reorientation, we increased BB disorientation by incubating *DISA-1* rescue cells at high ciliary forces (37°C for 24 h) and then initiated BB reorientation by acute DisAp expression at low ciliary forces (*DISA-1* rescue; 25°C; Fig. 1 A). BB orientation is defined by the axis of the BB and their associated SFs relative to the cell's anterior-posterior polarity (Fig. 1 B; Galati et al., 2014). This was monitored along with SF length relative to time after expression of WT DisAp. SF elongation begins within 8 h after DisAp expression, and BBs begin to reorient by 48 h. Thus, SF elongation initiates before BB reorientation (Fig. 1, Ci and Cii). Although mild recoveries in SF length and BB orientation were observed in the negative control (uninduced condition), they were attributed to leaky GFP-DisAp expression (Fig. S1 E). Importantly, SF length positively corresponds with proper BB orientation (Fig. 1 Ciii, single cell level  $R^2$  value = 0.87; and Fig. S1 D, individual BB level  $R^2$  value = 0.92). This further suggests that SF elongation is important for promoting BB orientation and reorientation.

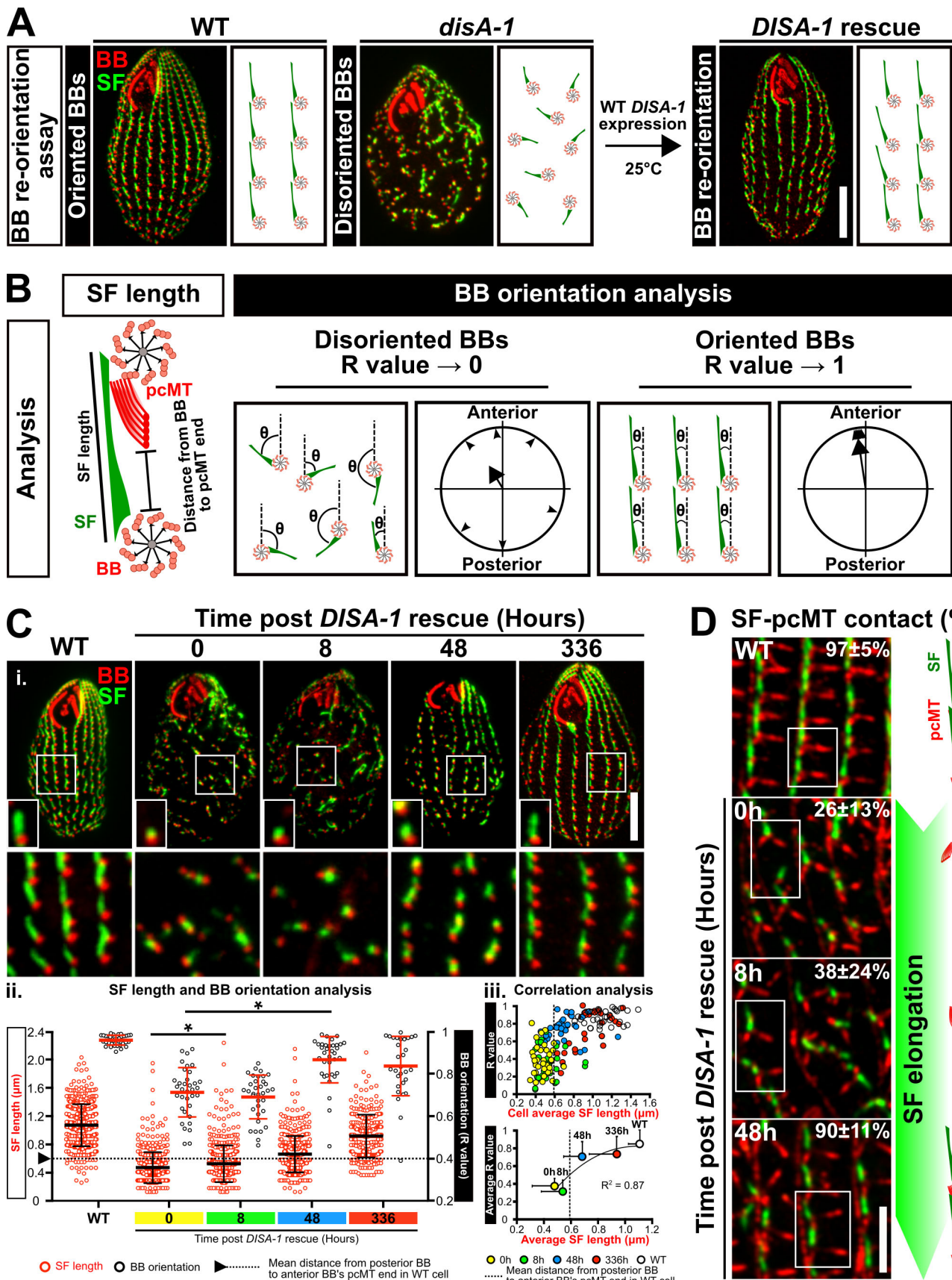
Since SF distal ends are normally juxtaposed to the anterior BB's pcMTs, we hypothesized that SF elongation in *DISA-1* rescue

cells enables disoriented BBs to establish attachments to the pcMTs of anterior BBs, thereby regaining BB orientation. To test this hypothesis, we measured the proportion of SF-pcMT interactions during *DISA-1* rescue. As SFs elongate in *DISA-1* rescue cells, the frequency of BBs with SF-pcMT interactions increases by 48 h (Fig. 1 D). Based on the relative position of the anterior BB's pcMT distal end in WT cells, SFs must attain a minimum length of  $0.59 \pm 0.25 \mu\text{m}$  to establish SF-pcMT interactions (Fig. S1 C). Consistent with this, we observed that disoriented BBs in *DISA-1* rescue cells began to regain BB orientation once the mean SF length surpassed this minimum length (Fig. 1, C and D; and Fig. S1 D).

To test whether BB reorientation occurs in single cells, we increased BB disorientation in *DISA-1* rescue cells using the same BB-disorientation regimen described earlier (high ciliary forces, 37°C for 24 h) and followed the rescue of single-cell *DISA-1* rescue isolates grown under cycling conditions. Upon the induction of *DISA-1* rescue at low ciliary forces (25°C), *DISA-1* rescue isolates underwent BB reorientation (Fig. S1 I, top). Interestingly, *DISA-1* rescue isolates recovered at different rates (Fig. S1 I, bottom). Since a subpopulation of cells still retains poor BB orientation even at 336 h after *DISA-1* rescue induction, we postulate that the difference in recovery rate results from the varying degree of BB disorientation at the onset of the experiment (Fig. 1 Ciii and Fig. S1 D). Cells with more severe BB disorientation may require a longer recovery time. Collectively, this suggests that *T. thermophila* cells possess error correction mechanisms to resolve BB disorientation. Indeed, spontaneous BB disorientation was observed in 5% of WT cells (Fig. S1 A). While SF-independent mechanisms, such as the potential role of neighboring cortical structures that promote BB reorientation, cannot be ruled out, we propose that SF elongation ensures the propagation of orientated BBs to future cell progeny.

In cycling cells, new BBs assemble, thereby reducing the spacing between neighboring BBs. This could promote BB reorientation during *DISA-1* rescue. To test this, cells were starved to inhibit both cell cycle progression and new BB synthesis. We first increased BB disorientation in *DISA-1* rescue cells before inhibiting cell cycle progression and BB duplication during *DISA-1* rescue. Non-cycling *DISA-1* rescue cells were equally efficient as cycling cells in BB reorientation. This suggests that cell cycle progression and new BB synthesis are not required for BB reorientation (Fig. S1, F and G). We postulate that this is because new BBs assemble from disoriented BBs, thereby further propagating BB disorientation.

We next tested whether ciliary beating promotes BB reorientation by translocating BBs to their proper position and orientation. To test whether ciliary beating is required for BB reorientation, we reduced ciliary beating during *DISA-1* rescue.  $\text{NiCl}_2$  treatment was used to inhibit dynein motor activity and ciliary beating (Larsen and Satir, 1991). In  $\text{NiCl}_2$ -treated media, disoriented BBs failed to reorient with reduced ciliary beating 48 h after *DISA-1* rescue (Fig. S1 H). However, SF length remained short, suggesting that cilia-dependent forces are necessary for SF elongation during *DISA-1* rescue. Because SFs did not elongate, we did not determine whether ciliary beating is important for promoting BB reorientation toward the cell's anterior pole.



**Figure 1. SFs promote BB reorientation.** (A) The BB reorientation assay. BB disorientation was exacerbated by shifting *DISA-1* rescue cells from 25°C to 37°C for 24 h. To promote BB reorientation, cells were shifted from 37°C to 25°C coincident with WT DisAp protein expression. (B) Schematic of SF length

measurements, distance measurements from the posterior BB to the anterior BB's pcMT distal end to establish the minimal SF length that is required for SF–pcMT contact, and BB orientation analyses. BBs and pcMTs, red; SF, green. (C) i and ii: DisAp expression leads to SF elongation before BB reorientation. BB reorientation occurs when SF length surpasses the minimal length that is required for SF–pcMT interactions (arrowhead and dotted line marks the mean distance from the posterior BB to the anterior BB's pcMT distal end in WT cells). SF length and BB orientation partially recover 2 wk after *DISA-1* rescue. BB, red; SF, green. All small insets show a representative BB and SF. i, bottom: White box marks region of interest (large inset). iii, top: Average SF length positively corresponds with BB orientation (R value) on the single cell level during *DISA-1* rescue. iii, bottom: Correlation analysis between the average SF length and BB orientation during *DISA-1* rescue. BB orientation in most cells is recovered once SF length surpasses the minimal length that is required to establish SF–pcMT contacts (dotted line marks the mean distance from the posterior BB to the anterior BB's pcMT distal end in WT cells). Data points are fitted with a polynomial (order = 2) function.  $n \geq 300$  SFs ( $\geq 30$  cells). Mann-Whitney test. \* denotes P value  $<0.01$ . Mean  $\pm$  SD. Scale bars, 10  $\mu\text{m}$  (cell), 1.3  $\mu\text{m}$  (small inset width), 7.8  $\mu\text{m}$  (large inset width). (D) Frequency of SF–pcMT contacts is increased as SF length recovers in *DISA-1* rescue cells. Schematic illustrates the position and orientation of two BBs within a region of interest (white box). BB-associated microtubule appendages, red; SF, green.  $n = 240$  SFs (24 cells). Scale bar, 5  $\mu\text{m}$ . tMT, transverse microtubule.

### Ciliary forces tune SF length

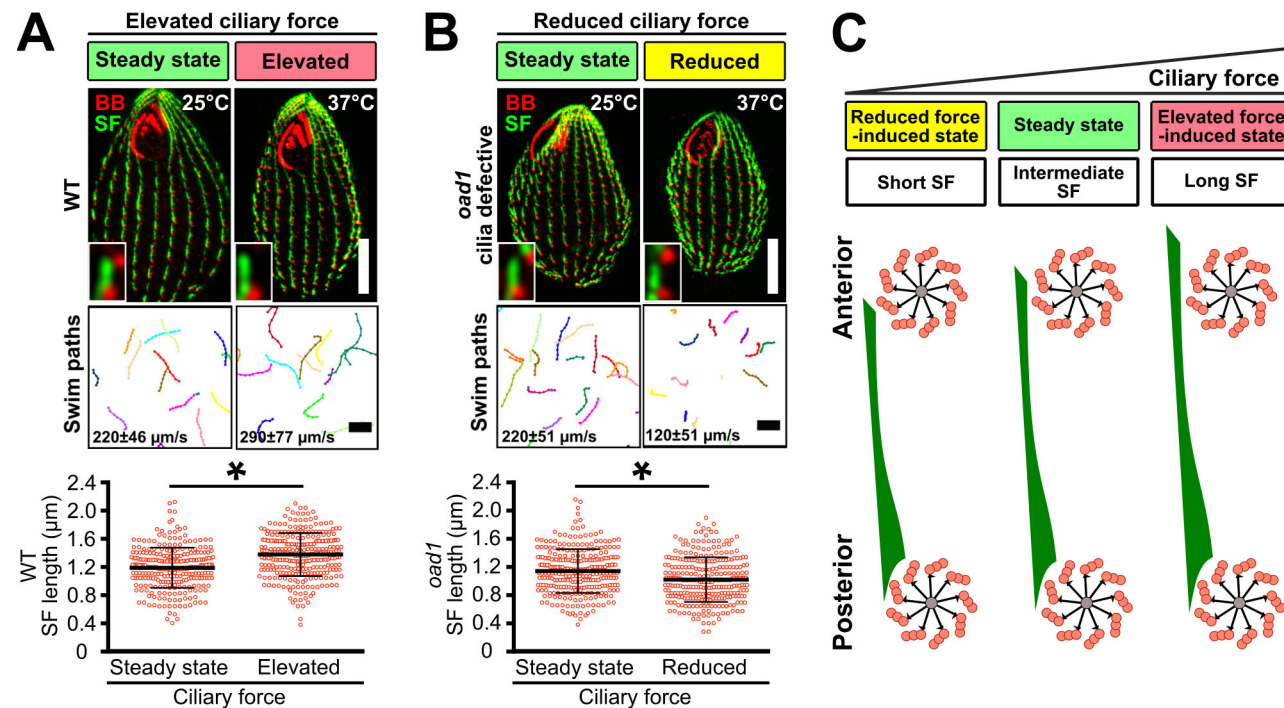
To investigate the relationship between ciliary forces and SF length, we exposed cells to a range of ciliary forces and quantified SF length. Consistent with our prior work, SFs of WT cells elongate by 16% when cell swimming and cilia-dependent forces are increased with elevated temperature for 4 h (elevated force-induced state; Fig. 2, A and C; Galati et al., 2014). Because SFs commonly overlap with each other (66%), the number of SFs that could be spatially resolved by conventional fluorescence microscopy was limited. Using structured illumination microscopy (SIM), a larger proportion of SF lengths was resolvable, and was quantified to show that SFs generally elongate with elevated ciliary forces (Fig. 2 A and Fig. S1 B).

SFs were previously shown to shorten at low temperatures (Galati et al., 2014). To establish whether SF length responds to

reduced ciliary forces independent of temperature effects, SF length was measured when ciliary beating was inhibited. To inhibit ciliary beating, the temperature-sensitive mutant strain, *outer arm deficient 1* (*oadl*; cilia lacking outer arm dynein), was grown at restrictive temperature (37°C for 4 h; Attwell et al., 1992). Elevated temperature reduces swimming rates of *oadl* cells by 45% and SF length by 11% (Fig. 2 B). Similarly, 6 h of NiCl<sub>2</sub> treatment reduces cell motility and SF length by 67% and 15%, respectively (Fig. S2 A). Thus, SF length is dynamically responsive to elevated and reduced ciliary forces (Fig. 2 C).

### SFs contact pcMTs and the cell cortex

Since SF elongation maintains BB orientation in WT cells and SFs promote BB reorientation in *DISA-1* rescue cells (Fig. 1; Galati et al., 2014), we postulated that SF length above a certain



**Figure 2. SFs elongate and shorten with ciliary force.** (A) Cell motility and SF length of WT cells were elevated by increasing temperature (37°C for 4 h). (B) Cell motility and SF length of the temperature-sensitive mutant strain, *outer arm deficient 1* (*oadl*), were reduced at restrictive temperature (37°C for 4 h; Attwell et al., 1992). BB, red; SF, green. All insets show a representative BB and SF. SF length quantitation:  $n = 300$  SFs (30 cells); motility assay, 90 cells. Mann-Whitney test. \* denotes P value  $<0.01$ . Mean  $\pm$  SD. Scale bars, 10  $\mu\text{m}$  (cell), 1.3  $\mu\text{m}$  (inset width), 100  $\mu\text{m}$  (swim paths). (C) Schematic illustrating SF length response to varying levels of ciliary force. BB-associated microtubule appendages are not shown.

threshold promotes SF interactions with structures to facilitate normal BB organization and orientation. Prior work revealed the close proximity and interactions of SFs to BB microtubule appendages and the cell cortex (Iftode et al., 1996; Pitelka, 1961). However, it is not known whether SF interactions persist at different ciliary forces. Here, we investigate SF interactions that promote BB positioning and orientation (Fig. 3 A).

To determine whether SFs interact with the pcMTs of anterior BBs at varying levels of ciliary force, we measured interactions between these structures using EM tomography of WT cells grown at normal (25°C, steady-state) or high ciliary force (38°C, elevated force-induced state). SFs extend anteriorly, crossing the surface of the anterior BBs' pcMT bundles in close proximity (Fig. 3 A*i*). We observed electron densities that link SFs and pcMTs, suggesting that BBs interact with neighboring BBs by forming bridges between SFs and pcMTs (Fig. 3 B and Video 1). These linkages were observed at both normal and high ciliary forces, indicating that, under all conditions measured, SFs physically link neighboring BBs via pcMTs (Fig. 3 B and Fig. S2 B). The SF-pcMT linkage length is longer in the elevated force-induced state (steady-state,  $16.0 \pm 4.6$  nm; elevated force-induced state,  $20.0 \pm 7.4$  nm; mean  $\pm$  SD;  $P$  value = 0.03). Because these linkages were observed under both steady-state and elevated force-induced state, they may be constitutive components required for the preservation of BB organization and orientation (Fig. S2 B).

Upon elevated ciliary-beating forces, SFs elongate beyond the anterior BB's pcMTs. This suggests that the SF distal ends may provide a secondary reinforcing interaction to resist elevated forces from ciliary beating. SFs are oriented toward the cell cortex, suggesting that SFs may have an interacting structure there (Fig. 3 A*ii*; Allen, 1967; Galati et al., 2014). To quantify the position of SFs relative to the cell cortex, we colocalized SFs with the cortical epiplasm protein EpiCp, fused to mCherry (Williams et al., 1987). Using a semiautomated image analysis routine, the distance between the peak intensities of SFs and EpiCp was quantified. SFs were not detectably closer to the epiplasm upon elongation (Fig. 3, C*ii* and C*iv*). However, SFs extended further along the epiplasm during the elevated force-induced state as compared with steady-state (Fig. 3, C*iii* and C*iv*).

To test for SF-cell cortex interactions, we employed EM tomography to monitor SF's distal end relative to the cortical epiplasm. Consistent with the above fluorescence quantification, SF-epiplasm interactions by longitudinal sections were not observed (Fig. 3 D and Fig. S2 C). However, we did observe electron-dense linkages between elongated SFs and the epiplasm surrounding the ciliary pocket in the elevated force-induced state, but not the steady-state (Fig. 3 E and Videos 2 and 3). While such interactions were not observed at steady-state, we cannot rule out the possibility that they are transient or were not captured. The length of the SF-epiplasm linkage was  $22.0 \pm 6.0$  nm (mean  $\pm$  SD). Unlike SF-pcMT connections, SF-epiplasm linkages were not observed at all SF-epiplasm interfaces around the ciliary pocket. SF-epiplasm linkages were detected in two of four SF-epiplasm interface tomograms during the elevated force-induced state. This is potentially due to transient interactions between these structures. Thus, SF-epiplasm linkages

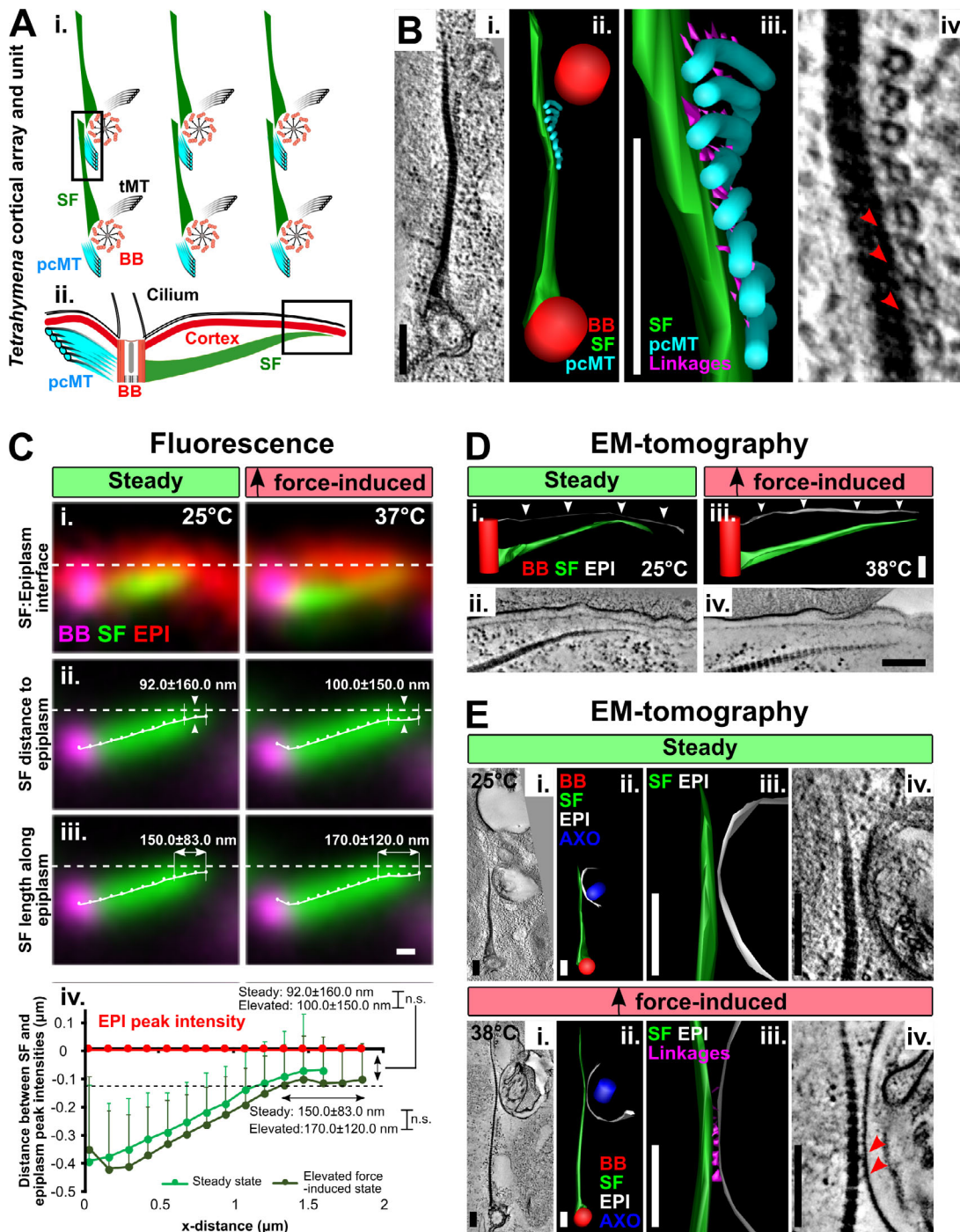
serve as dynamic, secondary interaction sites to reinforce BB organization and orientation when cilia and BBs experience greater mechanical forces.

### SFs are composed of a family of uniquely localized SF-assemblin (SFA) components

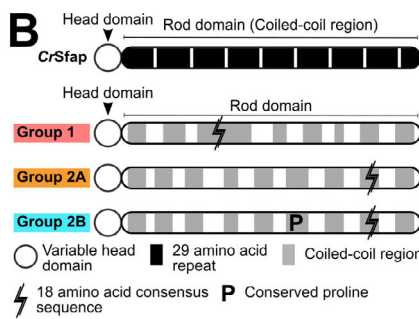
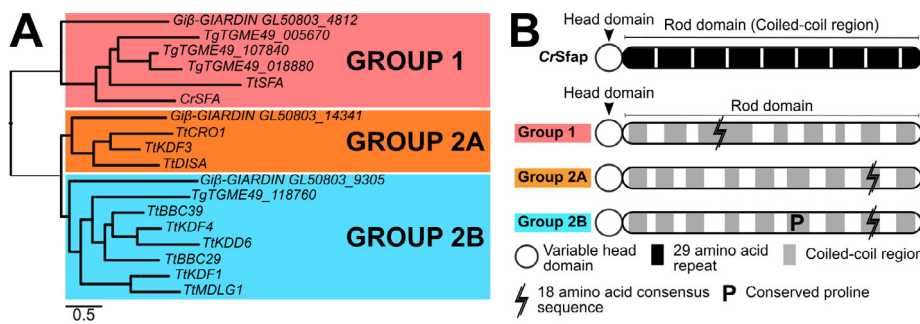
To determine the molecular composition of SFs, we performed phylogenetic analysis on the known SF gene *DISA-1* to identify similar genes. As *DisAp* belongs to the SFA family of proteins, we searched for SFA proteins across a broad set of eukaryotic species and found that SFA homologues are found exclusively among protists and Diphoda algae (Fig. 4 A and Figs. S4, S5, and S6; Galati et al., 2014). Moreover, these organisms possess variable numbers of SFA homologues, possibly resulting from whole-genome duplication and/or gene duplication events (Figs. S4, S5, and S6; number of SFA homologues: *Chlamydomonas reinhardtii*, 1; *T. thermophila*, 10; and *Paramecium tetraurelia*, 72). *T. thermophila* SFA homologues fall into three orthologous clades, which we designate Group 1, Group 2A, and Group 2B. Group 1 and Group 2 SFA proteins can be distinguished by the relative positions of an 18-amino acid consensus sequence (Fig. 4 B). This sequence appears near the N-terminus of Group 1 SFA proteins, but it is found on the C-terminus in Group 2 (Fig. 4 B and Fig. S2 E). In addition, Group 2B SFA proteins possess a single, conserved proline residue that distinguishes them from the Group 2A proteins (Fig. 4 B). Hence, we hypothesized that *T. thermophila* SFA proteins diverged and sub-functionalized.

To investigate whether *T. thermophila* SFA proteins localize to SFs, we fluorescently tagged these proteins in *T. thermophila* cells. From our localization analysis, we discovered that the Group 1 SFA homologue (Entrez Gene accession no. TTHERM\_00263290) localizes to cilia (Fig. S2 D), which is surprising given that its green algae homologue localizes to SFs (Lechtreck and Melkonian, 1991). Group 2 SFA proteins (Crolp, DisAp, Kdf3p, Kdd6p, Bbc39p, Kdf4p, Bbc29p, and Kdflp) localize to SFs (Galati et al., 2014; Chalker, D., personal communication). Using endogenously tagged genes under native promoters, we showed with SIM imaging that 8 out of 10 of the SFA proteins distinctly localize within SFs (Fig. 4 C). Thus, *T. thermophila* SF proteins exhibit unique phylogenetic and localization profiles.

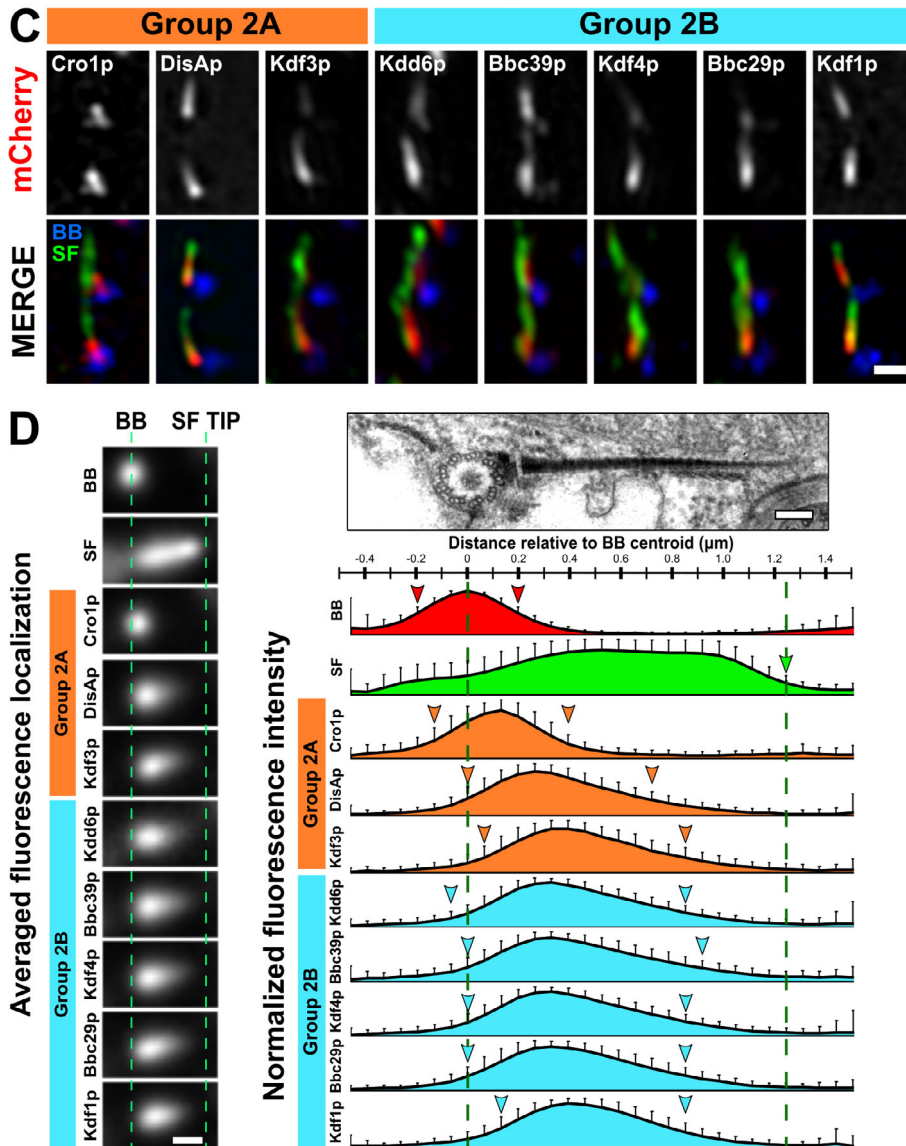
We developed a semiautomated, image-averaging pipeline to quantify the localization pattern for each *T. thermophila* SF protein (Fig. 4 D). Using BBs and SFs as fiducial marks, fluorescent images of each SF protein were aligned and averaged. To quantify the relative position and distribution of each SF protein along the SF, the fluorescent signal was measured. SF proteins localize to unique domains along the SF length (Fig. 4 D, left; and Fig. S2 F). Among Group 2A proteins, Crolp and DisAp localize to the SF base, while Kdf3p localizes more distally. Group 2B SF proteins (Kdd6p, Bbc39p, Kdf4p, Bbc29p, and Kdflp) localize along the SF length (Fig. 4 D, right). Because these components do not localize along the entire SF length, additional proteins likely form the SF distal ends. Coupled with the unique sequence features in each *T. thermophila* SFA group, it will be interesting to explore how these proteins influence SF assembly, length regulation, and function.



**Figure 3. SFs contact the pcMTs and the cell cortex.** **(A)** **i.** Schematic of *T. thermophila* cortical array (cross-sectional interface) illustrating BBs (red) and their associated appendage structures, including SFs (green), pcMTs (cyan) and transverse microtubules (tMT; black). **(ii)** Schematic depicting the SF-cell cortex longitudinal interface of a single *T. thermophila* cortical unit. Boxes mark the interfaces of interest. **(B)** EM tomographic images of the SF-pcMT interface between a pair of BBs at steady-state (25°C). Electron-densities (red arrowheads) link SFs to pcMT bundles (EM tomographic slice: **i** and **iv**; model: **ii** and **iii**). **(C)** **i.** Fluorescence images illustrating the SF position relative to the cell cortex (epiplasm; EpiC-mCherry [EPI]) at steady-state (25°C) and elevated ciliary force-induced state (37°C). **(ii and iii)** Averaged fluorescence images show that SFs extend toward (y axis) and along (x axis) the epiplasm at both steady-state and elevated force-induced state. BB, magenta; SF, green; EPI, red. Dashed line indicates the peak fluorescence intensities of EpiC-mCherry. **(iv)** Quantification of SF position relative to the epiplasm. Black dashed line marks the positions where the SF distal ends start to extend along the epiplasm. Steady-state,  $n = 104$  SFs (53 cells); elevated force-induced state,  $n = 94$  SFs (53 cells). Mann-Whitney test. Mean  $\pm$  SD. **(D)** EM tomographic images of the longitudinal SF-epiplasm interface. Linkages were not detected between SFs and the epiplasm (white arrowheads; EM tomographic slice: **ii** and **iv**; model: **i** and **iii**). **(E)** EM tomographic images of the cross sections of the SF-epiplasm interface at the ciliary pocket at steady-state (25°C) and elevated ciliary force-induced state (38°C). Electron-dense linkages (red arrowheads) were observed between SFs and the epiplasm only at the elevated force-induced state (EM tomographic slice: **i** and **iv**; model: **ii** and **iii**). B, D, E: BB, red; SF, green; pcMTs, cyan; epiplasm, white; linkages, magenta; axoneme (AXO), dark blue. Scale bars, 200 nm.



**Figure 4. SFs are composed of a family of uniquely localized SFA components.** (A) Simplified phylogenetic tree depicting the evolutionary relationships between SFA homologues of selected algae and protists. SFA homologues fall into three evolutionarily distinct groups. *Tt*: *T. thermophila*; *Cr*: *C. reinhardtii*; *Gi*: *G. intestinalis*; *Tg*: *T. gondii*. (B) Domain comparison of previously characterized green alga SFA protein (*C. reinhardtii*; Lechtreck and Silflow, 1997) and unique sequence features identified in SFA orthologous groups. (C) SIM imaging shows the unique localization of *T. thermophila* Group 2 SF proteins. BB, blue; SF, green; Group 2 SF proteins, white and red. Scale bar, 500 nm. (D) Left: Averaged fluorescence localization of *T. thermophila* Group 2 SF proteins. Number of SFs analyzed: *Cro1p*, 101; *DisAp*, 107; *Kdf3p*, 105; *Kdd6p*, 93; *Bbc39p*, 107; *Kdf4p*, 157; *Bbc29p*, 115; and *Kdf1p*, 109. SFs were gathered from  $\geq 60$  cells. Scale bar, 500 nm. Right: Schematic illustrating the relative localization of *T. thermophila* Group 2 SF proteins along the SF. BB, red; SF, green; Group 2A, orange; Group 2B, cyan. Green dashed lines mark the BB centroid and SF distal end. Arrowheads mark the start and end positions of the Group 2 SF proteins as defined by 30% from peak fluorescence intensity. SF start position is unmarked due to overlapping fluorescence signals from posterior SFs. Error bars indicate SD. Scale bar, 125 nm.



***Cro1p* promotes elevated ciliary force-induced SF elongation**  
 The localization of the ciliary row organizing-1 protein (*Cro1p*) to the SF base suggests that it may nucleate SFs, link SFs to BBs, and/or influence SF length (Fig. 4, C and D; Entrez Gene accession no. TTHERM\_000354599). The latter hypothesis is consistent with the proximal localizing SF component *DisAp*, which, when disrupted, causes SF shortening by ~60% (Fig. 1; Galati et al., 2014). To investigate the effect of *Cro1p* loss on SFs, *T. thermophila* *CRO1* was knocked out in the *T. thermophila*

macronucleus using CoDel (Hayashi and Mochizuki, 2015). Successful *CRO1* knockout was confirmed by the loss of *CRO1* genomic DNA and *CRO1* transcripts (Fig. S3, A–C). While exons 1–5 were eliminated, a partial transcript containing exons 6 and 7 was detected (Fig. S3 C, white box). Based on an alternate transcriptional start site within exon 7, expression of a truncated 20-amino acid *Cro1p* cannot be excluded. A *cro1Δ*+*CRO1* rescue strain was created by reintroducing *CRO1* into the *cro1Δ* cells (Fig. S3 D).

At steady-state, *cro1Δ* cells swim ~11% slower than WT cells (Fig. S3 E). At elevated forces, *cro1Δ* cells swim 60% slower than WT cells (Fig. S3 E). Thus, *cro1Δ*, like *disA-1* mutants, impedes cellular swimming behavior when ciliary beating forces are elevated.

To test whether the reduced cell motility in *cro1Δ* cells results from BB disorientation, we visualized BBs in *cro1Δ* cells. At steady-state, most BBs in *cro1Δ* cells were oriented. However, BB disorientation is increased at the elevated force-induced state (Fig. 5, A and B). Moreover, BB spacing is slightly increased in *cro1Δ* cells grown at elevated force-induced state as compared with WT cells, which retain normal BB spacing at both steady-state and elevated force-induced state (Fig. S3 K). Compared with *disA-1* mutants, *cro1Δ* cells exhibit intermediate BB disorientation (Fig. S3 L; Galati et al., 2014). We next tested whether *Cro1p* loss impacts SF length (Fig. 5 C). *cro1Δ* SFs are 18% shorter than WT SFs during steady-state and remain short at the elevated ciliary force-induced state (Fig. 5, A and C). Thus, *Cro1p* loss results in shorter SFs and an inability to lengthen SFs when ciliary forces are increased. Interestingly, the extent of the SF length defect in *cro1Δ* and *disA-1* cells correlates with the extent of BB disorientation (Fig. 5 D and Fig. S3 L; elevated force-induced state: WT,  $1.45 \pm 0.33 \mu\text{m}$ ; *cro1Δ*,  $0.90 \pm 0.35 \mu\text{m}$ ; *disA-1*,  $0.44 \pm 0.23 \mu\text{m}$ ; mean  $\pm$  SD). Unlike *disA-1* mutants, SFs in *cro1Δ* cells are long enough to establish SF–pcMT interactions and promote normal BB orientation at steady-state (Fig. 5, B and C; and Fig. S3 L). Hence, this further supports the importance of SF length in ensuring proper BB orientation (Fig. 5 D).

At steady-state, the majority of *cro1Δ* cells ( $76 \pm 9\%$ ; mean  $\pm$  SD) exhibit normal BB orientation. The remaining cells exhibit either individual BBs that are locally disoriented and disconnected from their neighboring BBs in ciliary rows (locally disoriented cells:  $20 \pm 7\%$ ; mean  $\pm$  SD) or connected BBs that are positioned within twisted BB rows relative to the cell's anterior-posterior axis (twisted cells:  $4 \pm 2\%$ ; mean  $\pm$  SD). The elevated force-induced state resulted in a larger proportion of *cro1Δ* cells with BB disorientation (Fig. 5, A and B; normal BB orientation,  $10 \pm 8\%$ ; BB disorientation,  $90 \pm 8\%$  [locally disoriented,  $66 \pm 8\%$ ; twisted cells,  $34 \pm 8\%$ ], mean  $\pm$  SD). This suggests that SF–pcMT interactions alone are insufficient to maintain BB orientation under the elevated force-induced state. Moreover, it reveals that proteins that localize to the SF base possess distinct SF functions, whereby *DisAp* is required for ensuring proper SF length to achieve SF–pcMT interactions and *Cro1p* is required for elevated force-induced SF elongation (Fig. 5 D and Fig. S3 L). Based on the distinct functions of *DisAp* and *Cro1p*, we hypothesized that the double knockout of *DISA-1* and *CRO1* will lead to cells that do not assemble SFs. However, we failed to obtain clones of the double knockout strain. This suggests that the loss of function of both proteins is synthetically lethal.

The twisted cells are reminiscent of the *twisty* and *scREWY* mutants in *T. thermophila* and *P. tetraurelia*, respectively (Fig. S3 F; Frankel, 2008; Jerka-Dziadosz et al., 1995; Whittle and Chen-Shan, 1972). Reduction of ciliary forces by shifting *cro1Δ* cells from 39°C to 25°C for 24 h rescued twisted BB rows back to WT configuration, indicating that twisted BB rows arise from elevated ciliary forces (Fig. S3 G). Because SFs are generally

oriented in *cro1Δ* cells with a few examples of local disorientation, we predicted that locally disoriented BBs possess shorter SFs that cause BBs to disconnect from their anterior neighbor. Indeed, locally disoriented BBs possess shorter SFs than BBs in twisted BB rows, which leads to BB disconnection (Fig. S3, H–J and L). Cells with twisted BB rows, on the other hand, possess SFs that are long enough to establish and maintain BB connections, but they are unable to elongate at the elevated force-induced state (Fig. S3, H–J and L). We postulate that this makes BB rows more susceptible to asymmetric ciliary forces and causes BB rows to deviate from the cell's anterior-posterior axis (Fig. 5 A and Fig. S3, F and L; Cheung and Jahn, 1976). Collectively, this suggests that *Cro1p* ensures proper SF lengthening to establish secondary SF–cell cortex interactions during elevated ciliary forces (Fig. 5 D).

## Conclusions

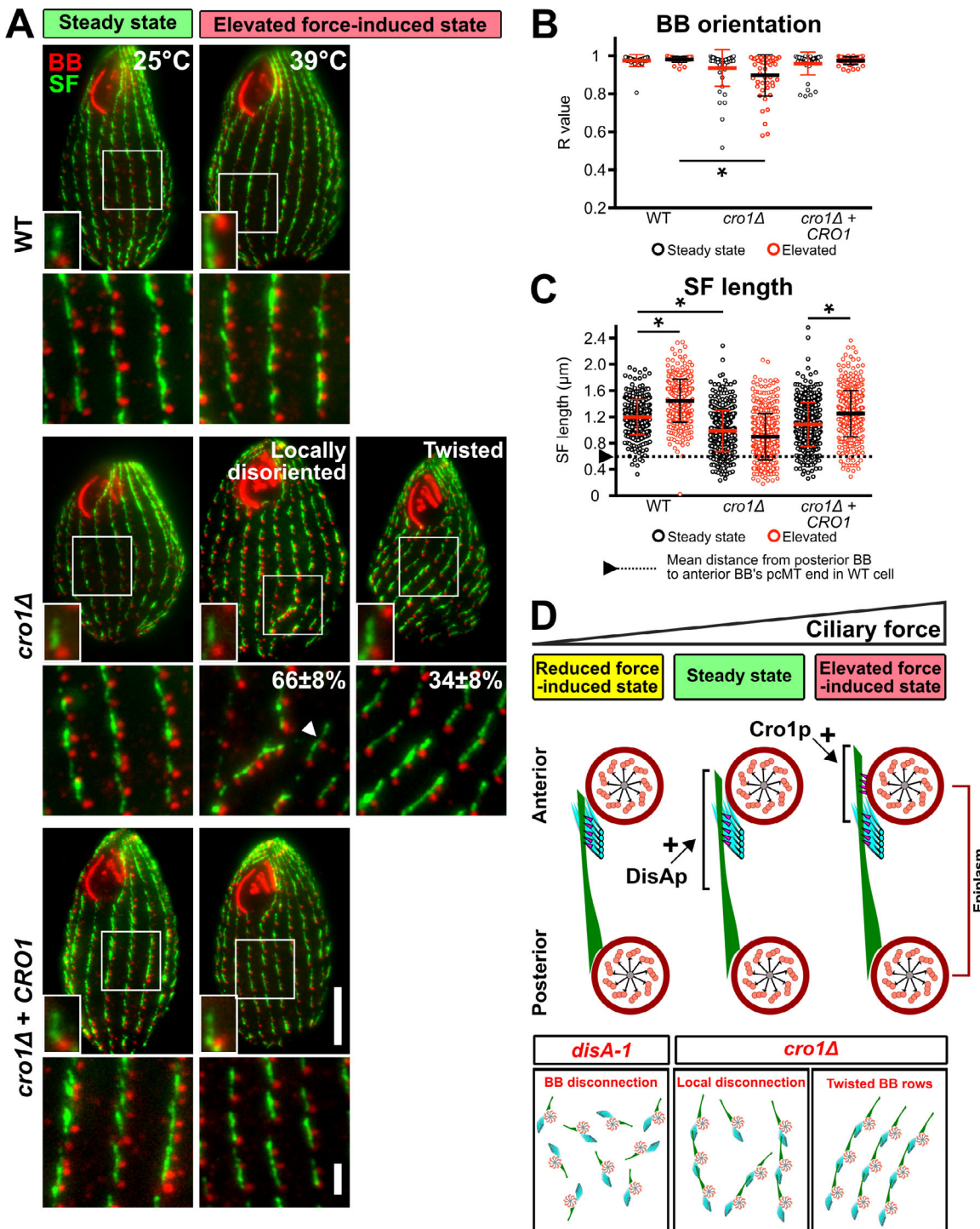
Here, we show that BB-associated SFs are ciliary force-responsive structures that attain unique length states to organize multiciliary arrays. To stabilize the positioning and orientation of BBs, SFs establish primary (steady-state) and secondary (elevated force-induced state) interactions with the anterior BB pcMTs and the cell cortex, respectively. This facilitates BB connections to propagate metachronal ciliary beating for cellular motility. Consistent with the sub-functionalization of SF components, we discovered that one proximally localized SF protein, *DisAp*, maintains steady-state SF lengths while a second proximal SF protein, *Cro1p*, is required for elevated force-induced SF elongation. We show that SF length maintenance and elongation promote attachments between neighboring BBs and to the cell cortex to ensure proper BB positioning and orientation.

## Materials and methods

### *T. thermophila* culture

*T. thermophila* cells were grown in 2% SPP media (2% proteose peptone, 0.2% glucose, 0.1% yeast extract, and 0.003% Fe-EDTA). For studies under cycling conditions, cells were analyzed at mid-log phase ( $4-5 \times 10^5$  cells/ml) as determined using a Coulter Counter Z1 (Beckman Coulter). For studies under non-cycling conditions, cells were arrested in the G1 phase of the cell cycle by washing and culturing in 10 mM Tris-HCl, pH 7.4, for 48 h. Cell concentration was kept at  $4-5 \times 10^5$  cells/ml. For microscopy experiments, analyses were restricted to nondividing cells as judged by those lacking a developing oral structure.

To study BB reorientation, *DISA-1* rescue cells were grown in SPP media at high ciliary forces (37°C for 24 h) to exacerbate BB disorientation. Next, cells were grown under cycling condition (SPP media) or noncycling condition (10 mM Tris-HCl, pH 7.4) at steady-state (25°C). Under both conditions, experiments were performed with and without GFP-*DisAp* expression. For the single cell rescue experiment, *DISA-1* rescue cells were subjected to the same BB disorientation regimen (high ciliary forces, 37°C for 24 h) before they were isolated and grown under cycling condition (SPP media) with and without GFP-*DisAp* expression. Cells were collected at the respective time points, fixed for immunofluorescence assay, and imaged. The assessment criteria



**Figure 5. *Cro1p* promotes elevated ciliary force-induced SF elongation.** **(A)** Representative images of WT, *cro1Δ*, and *cro1Δ* rescue cells during steady-state (25°C) and elevated force-induced state (39°C for 24 h). More *cro1Δ* cells exhibit local BB disorientation (white arrowhead) and twisted BB rows compared with WT cells at elevated force-induced state. The percentage of each phenotype is calculated based on *cro1Δ* cells that exhibit BB disorientation. All small insets show a representative BB and SF. Scale bars, 10  $\mu$ m (cell), 1.3  $\mu$ m (inset width), 2  $\mu$ m (BB rows inset). **(B)** Quantification of BB orientation (R value) of WT, *cro1Δ*, and *cro1Δ* rescue cells. *cro1Δ* cells exhibit BB disorientation at elevated force-induced state. **(C)** Quantification of SF length of WT, *cro1Δ*, and *cro1Δ* rescue cells. SF length of *cro1Δ* cells failed to elongate at the elevated force-induced state. Arrowhead and dotted line mark the minimal SF length that is required for SF–pcMT interactions.  $n \geq 350$  SFs ( $\geq 40$  cells). Mann-Whitney test. \* denotes  $P < 0.01$ . Mean  $\pm$  SD. **(D)** Model depicting how SF base proteins, DisAp and Cro1p, influence SF length. Loss-of-function of these proteins either affect primary (SF–pcMT) or secondary (SF–cell cortex) interactions, and lead to varying degrees of BB disorganization and disorientation. A and D: BB and epiplasm, red; SF, green; pcMTs, cyan; linkages, magenta.

for the rescue of *DISA-1* rescue isolates are normal (i) cell morphology and (ii) swim path.

To expose cultures to reduced ciliary forces, *T. thermophila* cells were propagated in 2% SPP supplemented with 3 mM NiCl<sub>2</sub> (Sigma-Aldrich), which was added directly to the culture vessel from a 1 M stock. Dynein-dependent ciliary beating is inhibited by NiCl<sub>2</sub>, which blocks plasma membrane calcium channels and directly inhibits dynein motors (Larsen and Satir, 1991). For experiments that require an extended duration of NiCl<sub>2</sub> treatment, a lower dosage of NiCl<sub>2</sub> (2 mM) was used.

### Plasmids and *T. thermophila* strain construction

The BB reorientation assay was performed using the *DISA-1* rescue strain previously described (Galati et al., 2014). To initiate rescue in the *DISA-1* rescue strain, *DISA-1* expression was induced via a cadmium (II) chloride-driven promoter.

Previously reported *T. thermophila* strains overexpressing SFA proteins (Crolp, Kdf3p, Kdd6p, Bbc39p, Kdf4p, Bbc29p, and Kdflp) localize to the SF (www.suprdb.org; Chalker, D., personal communication). In this study, we created endogenously tagged genes under native promoters. WT *T. thermophila* cells (B1868) that express mCherry-tagged SF proteins were created using the p4T2-1-mCherryLAP construct (Winey et al., 2012). Briefly, the mCherryLAP cassette integrates at the endogenous target gene locus, and gene expression remains under the control of the endogenous promoter. Transformed cells were selected for paromomycin resistance and assayed to at least 2 mg/ml of paromomycin. For information on oligo design, please refer to Table S1.

### Phylogenetic analysis

We gathered an initial set of SFA-related protein sequences from our in-house orthology dataset by running Orthofinder 2.1.2 (Emms and Kelly, 2015) on a curated set of 169 eukaryotic proteomes with DIAMOND (Buchfink et al., 2015). To identify sequences that may have been missed, we constructed a custom hidden Markov model with the HMMER package (Eddy, 2011) using merged orthologous group sequences that were aligned with MAFFT (v7.271; Katoh and Standley, 2013). The final alignment was constructed by aligning the inclusive set of sequences with CulstalOmega (v1.2.1; Sievers et al., 2011), and curated by removing variable N- and C-termini sequences. The final set contains 205 SFA sequences (supplemental data file 1). Phylogenetic trees were constructed using RaxML (v8.2.4; Stamatakis, 2014) and MrBayes (v3.2.6; Ronquist et al., 2012). Both algorithms generated phylogenetic trees that are in agreement. Raw data are available on request. Since a suitable outgroup could not be determined, the final trees were rooted such that each branch contains eukaryotic-wide taxonomic distributions. Coiled-coil regions were predicted during manual curation of SFA protein sequence alignment (Fig. 4 B).

### Immunocytochemistry

#### BB and SF labeling

For immuno-cytochemical analyses of BBs and SFs,  $7 \times 10^5$  cells were pelleted at 600 g in a 1.5-ml microcentrifuge tube and fixed for 20 min with 1 ml of 70% ethanol + 0.1% Triton X-100 on ice. Cells were washed thrice with 10 mM PBS, pH 7.4, and blocked

in 0.5% BSA/PBS for 1 h at 4°C. Cells were immunostained by incubating in primary antibody (mouse anti-SF [5D8], 1:500; rabbit anti-centrin [BB], 1:500; Jerka-Dziadosz et al., 1995; Stemm-Wolf et al., 2005) overnight at 4°C, followed by secondary antibody (goat anti-mouse Alexa Fluor 488 or 594, 1:2,000; goat anti-rabbit Alexa Fluor 647, 1:2,000; Invitrogen) for a 2-h incubation at 4°C. Cells were mounted in Citifluor mounting media using #1.5 coverslips and sealed with nail polish. All antibodies were diluted in 0.5% BSA/PBS. Cells were washed thrice with 0.5% BSA/PBS after primary and secondary antibody incubations.

#### SF and pcMT labeling

For immuno-cytochemical analyses of SFs and pcMTs,  $7 \times 10^5$  cells were pelleted at 600 g in a 1.5-ml microcentrifuge tube and fixed for 5 min with 1 ml of 3.2% PFA/PHEM + 0.24% Triton X-100 (PHEM, 60 mM Pipes, 25 mM Hepes, 10 mM EGTA, and 2 mM MgCl<sub>2</sub>, pH 6.9) at 25°C. Next, cells were pelleted and re-suspended in 1 ml of ice-cold 0.5% Triton X-100 (PHEM) for 10 min on ice at 4°C. Cells were washed thrice with PHEM and blocked in 0.5% BSA/PHEM for 1 h at 4°C. Cells were immunostained by incubating in primary antibody (mouse anti-SF [5D8], 1:500; rabbit anti-acetylated α-tubulin lysine 40 [D20G3; Cell Signaling Technology], 1:100) overnight at 4°C followed by secondary antibody (goat anti-mouse Alexa Fluor 488 or 647, 1:2,000; goat anti-rabbit Alexa Fluor 594, 1:2,000; Invitrogen) for a 2-h incubation at 4°C. Cells were mounted in Citifluor mounting media using #1.5 coverslips and sealed with nail polish. All antibodies were diluted in 0.5% BSA/PHEM. Cells were washed thrice with 0.5% BSA/PHEM after primary and secondary antibody incubations.

### Light microscopy

Imaging experiments (Figs. 3 C, 4 D, 5 A; S1, A and F–I; S2 D, and S3, D and F) were performed with an inverted widefield microscope (Ti Eclipse; Nikon). A 100× Plan-Apochromat (NA 1.4) objective lens (Nikon) was used. Images were captured with a z-step size of 300 nm using a scientific complementary metal-oxide semiconductor camera (Zyla; Andor Technology).

Super-resolution localization experiments of SF-pcMT interactions (Figs. S1 C and S3 L) and Group 2 SF proteins (Fig. 4 C) were performed via SIM with the Nikon 3D SIM system (Ti 2 Eclipse). A 100× total internal reflection fluorescence objective (NA 1.45) was used. Images were captured with a complementary metal-oxide semiconductor camera (Orca-Flash 4.0; Hamamatsu) with a z-step size of 300 nm. Raw SIM images were reconstructed by the image stack reconstruction algorithm (Nikon Elements).

Confocal microscopy was performed using an inverted microscope (Ti Eclipse) with a 100× Plan-Apochromat (NA 1.43) objective lens (Nikon) and a swept field confocal scan head with the 35-μm slit mode (Prairie Technologies). Images were captured with a charge-coupled device camera (iXon X3; Andor Technology). Confocal images were also acquired with the A1 confocal laser microscope (Nikon). All images were acquired with Nikon Elements with a z-step size of 300 nm at room temperature (Figs. 1, 2, S1 E, and S2 A).

## Electron tomography

Cells were prepared for electron tomography as previously described (Giddings et al., 2010; Meehl et al., 2009). Cells were gently spun into 15% dextran (molecular weight 9,000–11,000 D; Sigma-Aldrich) with 5% BSA in 2% SPP. A small volume of concentrated cells was transferred to a sample holder and high-pressure frozen using a Wohlwend Compact 02 High Pressure Freezer (TechnoTrade International). After low-temperature freeze substitution in 0.25% glutaraldehyde and 0.1% uranyl acetate in acetone, cells were slowly infiltrated with Lowicryl HM20 resin. Serial thick (250–300 nm) sections were cut using a Leica UCT ultramicrotome. The serial sections were collected on Formvar-coated copper slot grids and poststained with aqueous uranyl acetate followed by Reynold's lead citrate.

Dual-axis tilt series ( $-60$  to  $+60^\circ$ ) of *T. thermophila* cells were collected on a FEI Tecnai 300kV FEG-TEM (FEI). Images were acquired using the SerialEM acquisition program (Mastronarde, 2005) with a Gatan OneView (4k  $\times$  4k) camera. Serial section tomograms of *T. thermophila* cortical structures were generated using the IMOD 4.9 software package (Kremer et al., 1996; Mastronarde, 1997). Three tomograms for each interface of interest were analyzed.

## *T. thermophila* motility measurements

*T. thermophila* cell motility was imaged on an inverted widefield microscope using a 20 $\times$  objective lens (Ti Eclipse; Nikon). Each movie duration is 2.5 s (exposure duration, 50 ms; frame rate, 20 frames per second). To quantify swim rates, the relative displacement of the anterior or posterior pole of cells was tracked for 500 ms via the FIJI MTrackJ plugin (Meijering et al., 2012). Analyses were restricted to cells that swam along the same x y plane.

## Image analysis: SF length, BB orientation (R value), distance between posterior BB and anterior BB's pcMT distal end, and frequency of SF-pcMT contact

SF length and BB orientation were quantified by a semi-automated strategy previously described (Galati et al., 2014). Briefly, all images were first uniformly contrasted before quantitation. SF length was measured via the freehand tool (Fig. 1 B). To compute BB orientation (R value), which reflects the variance of BBs' orientation relative to the cell's anterior-posterior axis, the ORIANA circular statistics suite (Kovach Computing Services) was used. Based on the displacement of the SF from its associated BB and the anterior pole of the cell, an angular measurement was obtained (Fig. 1 B). For each *T. thermophila* cell, at least five angular measurements were gathered from SFs that are positioned within a 10- $\mu$ m box placed at the cell's medial region. Using the angular measurements, the mean vector and length of the mean vector (R value) were calculated. The BB orientation in each cell is represented by a single R value. An R value of 1 indicates that SFs are uniformly oriented toward the anterior pole of the cell. Conversely, an R value of 0 indicates that SFs are randomly oriented. The discrepancy in R values between our prior publication and this paper is attributed to slight differences in SF sampling between experimentalists (Galati et al., 2014). Consistency of R-value measurements was

ensured in this paper. Cell level SF length:BB orientation correlation analysis was performed by correlating the average SF length within the same cell to its corresponding R value. Individual BB level SF length:BB orientation correlation analysis was performed by correlating the SF length of an individual BB to its SF's angular displacement from the anterior pole of the cell.

To quantitate the position of the anterior BB's pcMT distal end (relative to the posterior BB), we colocalized SFs with pcMTs and measured the distance from the pcMT distal end to the posterior BB (determined by the position of SF base; Fig. 1 B). To measure the frequency of SF-pcMT contacts, we determined the criteria for contact to be an overlap of the SF and pcMT fluorescence signals by at least 220 nm (2 pixels). Both analyses were restricted to SFs and pcMTs found at the cell's medial region.

## Image analysis: Quantification of SF position relative to the cell cortex (epiplasm)

Fluorescence quantification of SF position relative to the cell cortex was performed by a semiautomated strategy that utilizes the FIJI macro scripting language and plugins (Schindelin et al., 2012). Image stacks colocalized for BBs, SFs, and the cell cortex (marked by EpiC-mCherry) were preprocessed to generate maximum projections (three slices; z-step size = 300 nm [FIJI Z Project; Max Intensity plugin]). All analyses were restricted to BBs and their associated SFs that were found in the medial region of the cell. To visualize the longitudinal SF-cell cortex interface, BBs positioned at the side of *T. thermophila* cells were selected. Each BB unit, which includes a BB, its associated SF, and the nearby epiplasm, is cropped as an individual image (5.2  $\mu$ m  $\times$  5.2  $\mu$ m) for further processing. The average distance between the SF and the epiplasm is measured based on the peak intensities of the SF (distal 25th percentile of its total length; labeled with mouse anti-SF [5D8]) and the epiplasm marker EpiC-mCherry (FIJI Plot Profile plugin). The average SF length that spans along the epiplasm is quantified based on SF peak fluorescence intensities that fall within 130 nm (2 pixels) from the EpiC-mCherry peak fluorescence intensities.

## Image analysis: Fluorescence image averaging

Fluorescence image averaging was performed by a semi-automated strategy that utilizes the FIJI macro scripting language and plugins (Schindelin et al., 2012). Image stacks were preprocessed to generate maximum projections of the *T. thermophila* cell side that is nearer to the glass cover (11 slices; z-step size = 300 nm [FIJI Z Project; Max Intensity plugin]). All analyses were restricted to BBs and their associated SFs that were found in the medial region of the cell. Each selected BB and its associated SF (SF length: 1.1–1.3  $\mu$ m) is cropped as an individual image (5.2  $\mu$ m  $\times$  5.2  $\mu$ m) for further processing.

Next, selected BBs and SFs served as fiducial marks for 2D alignment. To align BBs, the brightest pixel within each BB is identified by Gaussian blur (FIJI Gaussian Blur plugin;  $\sigma$  = 2 pixels). Based on the position of the brightest pixel in each BB, BBs were aligned through the FIJI Translate plugin. Next, a second alignment step based on the SF distal end was performed. To align SFs along the same axis, SFs were rotated (relative to the BB's brightest pixel) using the FIJI Rotate plugin.

Prior to fluorescence image averaging, the local background fluorescence intensities (defined by the regions directly adjacent to the signal of interest) were measured, averaged, and subtracted from each respective channel. Next, all peak intensities for each respective channel were normalized to 1. Finally, fluorescence images of aligned BBs and SFs were averaged using the FIJI Average Intensity plugin.

To measure the average length distributions and relative start positions of Group 2 SF proteins along the SF, we measured the fluorescence distribution of each mCherry-tagged Group 2 SF protein (FIJI Plot Profile plugin). To quantify the length of SF protein-mCherry localization along the SF, a fluorescence-based criterion was used (intensity value  $\geq 30\%$  from the peak fluorescence intensity qualifies for length measurement). The start positions of Group 2 SF protein-mCherry along the SF are determined based on their relative positions from the BB centroid. The same fluorescence-based criterion was applied.

### Statistical analysis

All datasets were assessed for normal distribution using Shapiro-Wilk normality test. Student's *t* test was performed on normally distributed datasets. Mann-Whitney test was performed on datasets that do not conform to normal distribution. Tests for significance were unpaired and two-tailed. Categorical datasets were analyzed using  $\chi^2$  test. All error bars indicate SD. Statistical significance was set at *P* value  $<0.01$ . All analyses were performed on samples obtained from three independent experiments.

### Online supplemental material

Fig. S1 shows that SF elongation promotes BB reorientation. Fig. S2 shows SF length and position during varying levels of ciliary force and the molecular composition of SFs. Fig. S3 shows that Crolp promotes elevated ciliary force-induced SF elongation. Figs. S4, S5, and S6 show phylogenetic analysis of SFA homologues among ciliates and algae; rooted phylogenetic trees of 205 SFA protein sequences gathered from 50 species of protists and algae; and three taxonomically diverse clades are identified and designated as Group 1, Group 2A, and Group 2B. Figs. S4, S5, and S6 show Group 1, Group 2A and Group 2B, respectively. Video 1 shows serial, tomographic slices and model depicting cross section of SF-PCMT interface at steady-state (25°C). Video 2 demonstrates serial, tomographic slices and model depicting cross section of SF-epiplasm interface at steady-state (25°C). Video 3 shows serial, tomographic slices and model depicting cross section of SF-epiplasm interface at elevated force-induced state (38°C). Table S1 provides tabulation of oligo sequences used for fluorescent protein tagging and the functional study of *T. thermophila* Crolp. The supplemental data file 1 provides multiple sequence alignment of 205 SFA protein sequences from 50 protist and algae species.

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Author contributions: A.W.J. Soh designed and executed experiments and data analyses, and wrote the manuscript. T.J.P. van Dam performed the phylogenetic analysis. A.J. Stemm-Wolf generated *Tetrahymena* strains. A.T. Pham performed the DISA-1 rescue experiments. G.P. Morgan and E.T. O'Toole acquired EM tomograms. C.G. Pearson directed the study and wrote the manuscript.

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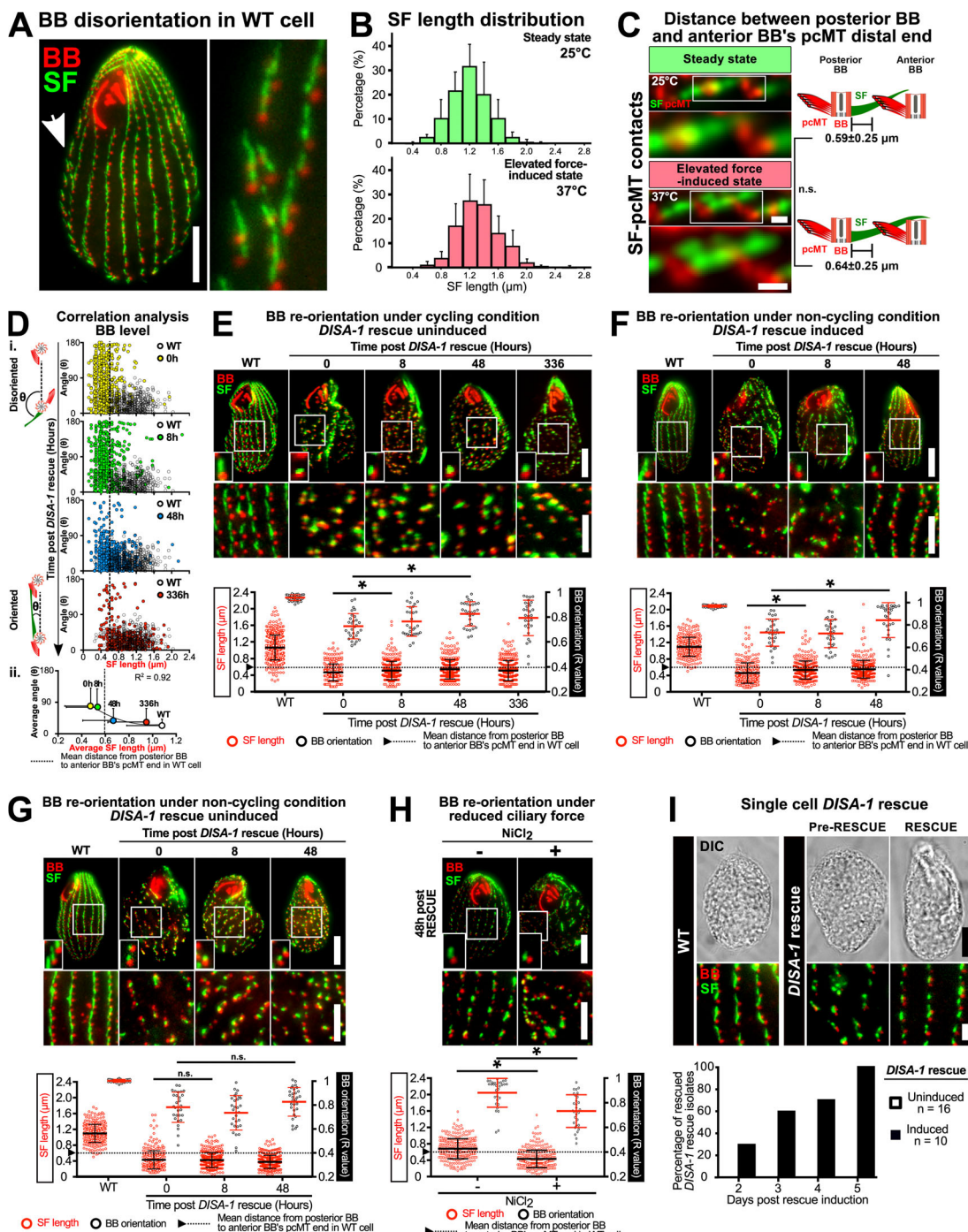
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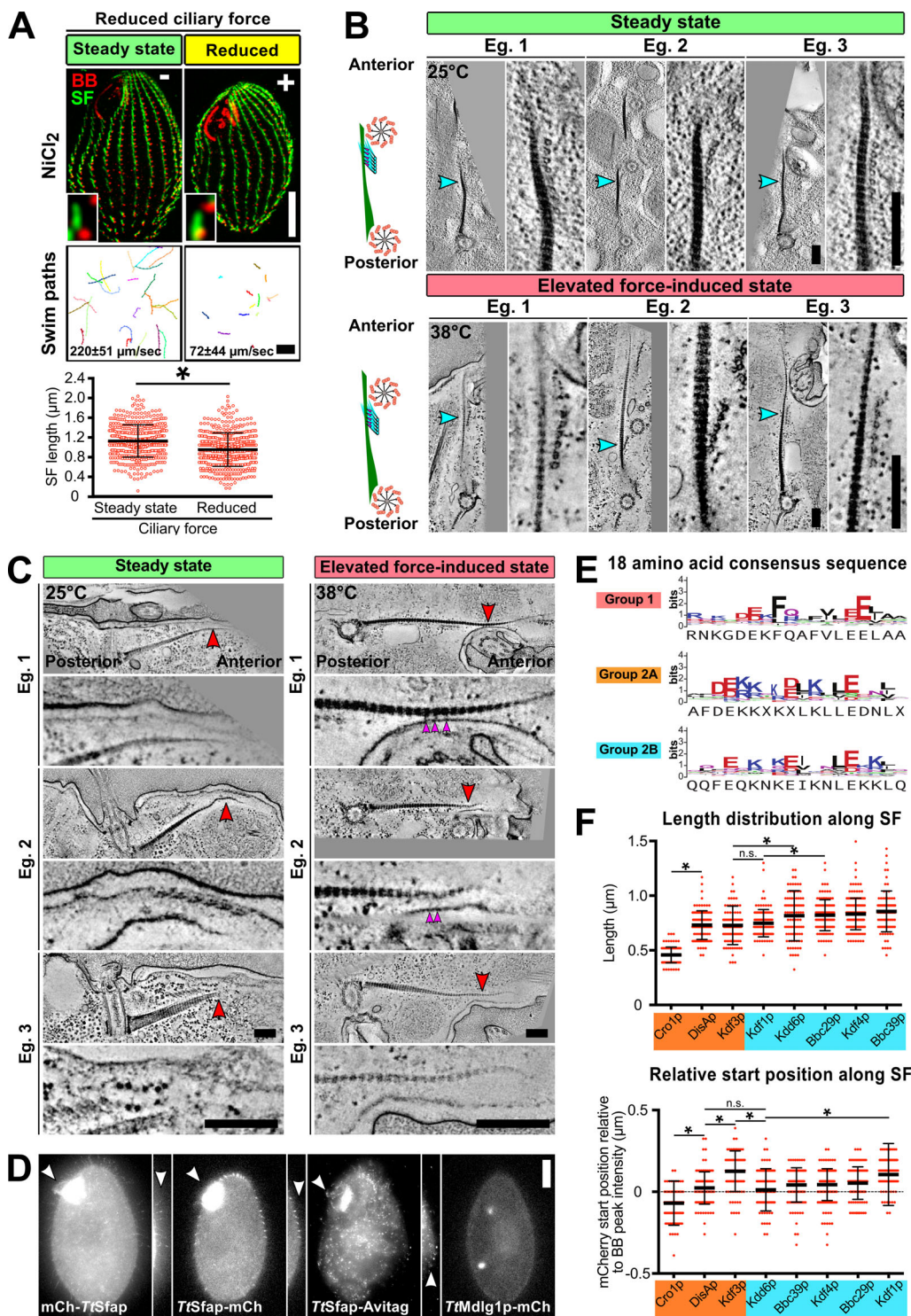
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## Supplemental material

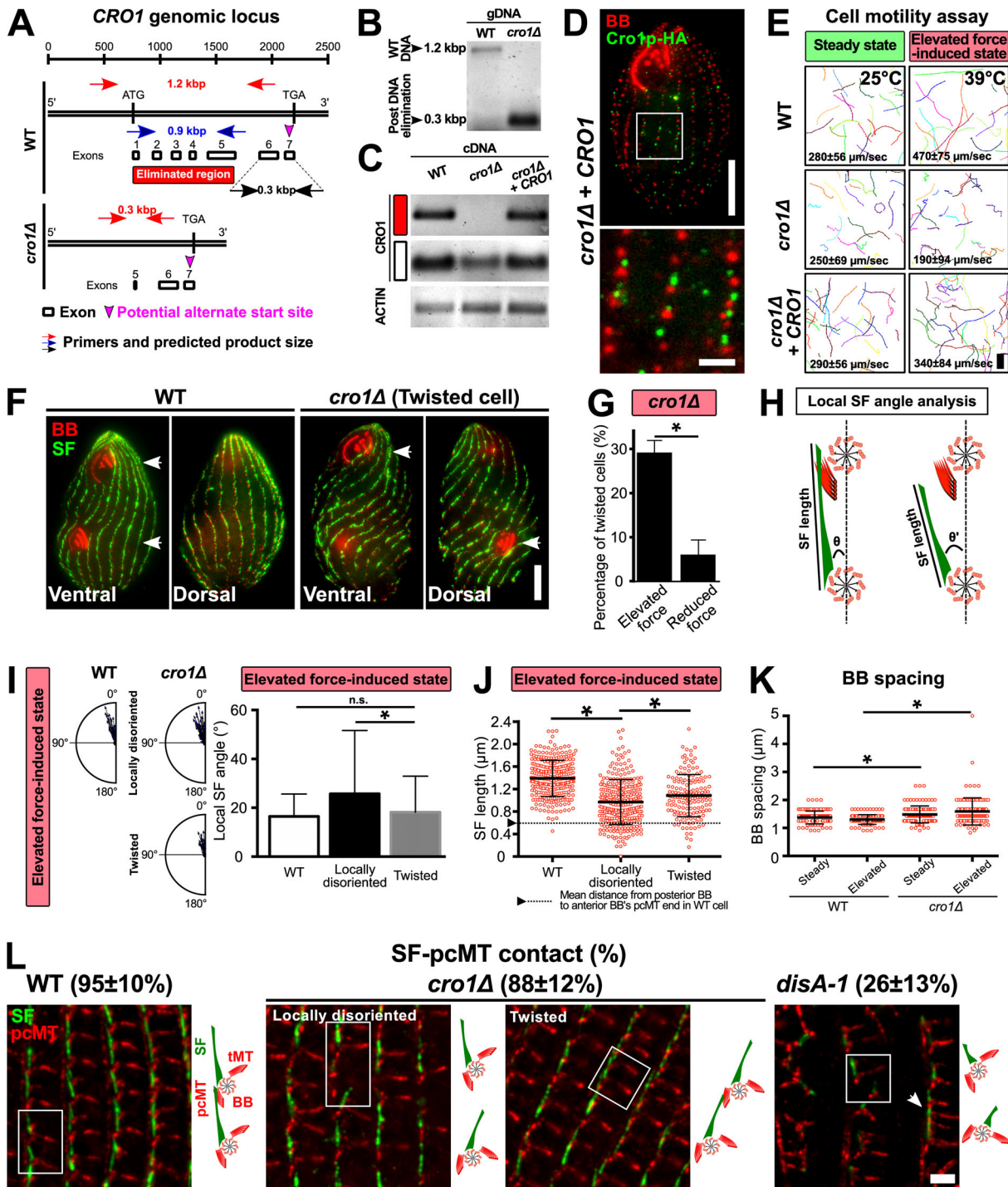
Soh et al., <https://doi.org/10.1083/jcb.201904091>



**Figure S1. SFs promote BB reorientation.** **(A)** Widefield image depicting BB disorientation in WT cell. Arrow marks disoriented BBs. Scale bar, 10 µm (cell), 4.5 µm (inset width). **(B)** Distribution of SF length measured from SIM-acquired SF-BB images. **(C)** Distance measurements between posterior BB and anterior BB's pcMT distal end in WT cells at steady-state and elevated force-induced state. Scale bars, 500 nm. **(D)** i: SF length positively corresponds with proper BB orientation (angle: SF angular displacement from its associated BB to the cell's anterior pole) on the individual BB level during DISA-1 rescue. ii) Correlation analysis between the average SF length and BB orientation (angle) during DISA-1 rescue. Most BBs regain proper orientation once the length of their SFs surpasses the minimal length to establish SF-pcMT contacts (dotted line marks the mean distance from the posterior BB to the anterior BB's pcMT distal end in WT cells). Data points are fitted with a polynomial (order = 2) function. **(E-H)** Negative control of BB reorientation analysis under cycling condition. Recoveries in SF length and BB orientation were attributed to leaky DisAp expression. **(F and G)** BB reorientation analysis under noncycling conditions. **(F)** Under noncycling conditions, DISA-1 rescue promotes SF lengthening before BB reorientation. **(G)** Negative control of BB reorientation analysis under noncycling condition. SF length and BB orientation recovery do not occur without DisAp expression. **(H)** BB reorientation analysis under reduced ciliary force. Under reduced ciliary force, SFs remained short and BBs failed to reorient. All small insets show a representative BB and SF. **(B-H)**  $n = \geq 300$  SFs ( $\geq 30$  cells). Mann-Whitney test. \* denotes  $P$  value  $< 0.01$ . Mean  $\pm$  SD. **(E-H)** Scale bars, 10 µm (cell), 1.3 µm (small inset width), 5 µm (BB rows inset). **(I)** Top: DISA-1 rescue isolates are rescued upon DisAp expression. Scale bars, 10 µm (cell), 2 µm (inset). **(I)** Bottom: DISA-1 rescue isolates recover at different rates. BB, red; SF, green.



**Figure S2. SF length and position during varying levels of ciliary force and the molecular composition of SFs.** (A) Cell motility and SF length of WT cells were reduced by  $\text{NiCl}_2$  treatment ( $25^\circ\text{C}$  for 6 h). BB, red; SF, green. Insets show a representative BB and SF. SF length quantitation:  $n = 500$  SFs (35 cells); motility assay, 90 cells. Scale bars, 10  $\mu\text{m}$  (cell), 1.3  $\mu\text{m}$  (inset width), 100  $\mu\text{m}$  (swim paths). (B) EM tomographic images of cross-sectional SF-PCMT interface at steady-state ( $25^\circ\text{C}$ ) and elevated ciliary force-induced state ( $38^\circ\text{C}$ ). Electron-dense linkages exist between SFs and PCMT bundles of the anterior BB (cyan arrowheads, enlarged interface). Scale bars, 200 nm. (C) EM tomographic images of longitudinal and cross-sectional SF-epiplasm interfaces at steady-state and elevated ciliary force-induced state (red arrowheads, enlarged interface; magenta arrowheads, SF-epiplasm linkages). Scale bars, 200 nm. (D) Localization of non-SF localizing *Tetrahymena* SFA proteins. Multiple labeling strategies indicate that *TtSfap* localizes to oral and cortical cilia (white arrowheads). *TtMdlg1p* is found in vacuoles. mCh: mCherry. Scale bars, 10  $\mu\text{m}$  (cell), 6.5  $\mu\text{m}$  (inset width). (E) Sequence logos of group-specific consensus sequences in SFA proteins (X, nonconserved residue). (F) Quantitation of Group 2 SF protein distribution along the SF. Top: Length distribution of Group 2 SF proteins. Bottom: Start positions of Group 2 SF proteins relative to BB peak intensity. Number of SFs analyzed: *Cro1p*, 101; *DisAp*, 107; *Kdf3p*, 105; *Kdd6p*, 93; *Bbc39p*, 107; *Kdf4p*, 157; *Bbc29p*, 115; and *Kdf1p*, 109. SFs were obtained from  $\geq 60$  cells. Mann-Whitney test. \* denotes  $P$  value  $< 0.01$ . Mean  $\pm$  SD.



**Figure S3. Cro1p promotes elevated ciliary force-induced SF elongation.** **(A)** Schematic illustrating the genomic locus of *CRO1* and the site that was targeted for DNA elimination (red box). **(B)** PCR assessment confirmed DNA elimination at the targeted site of *CRO1* genomic locus. **(C)** RT-PCR assessment confirmed the absence of *CRO1* transcript expression in *cro1Δ* cells (red box). A partial *CRO1* transcript was expressed downstream of the DNA-eliminated region (white box). **(D)** Expression and localization of Cro1p-HA in *cro1Δ* rescue strain. BB, red; Cro1p-HA, green. Scale bars, 10 μm (cell), 2 μm (inset). **(E)** Motility assay of WT, *cro1Δ*, and *cro1Δ* rescue cells at steady-state and elevated force-induced state. Scale bar, 100 μm. Motility assay, 90 cells. **(F)** Mispositioned oral structure in dividing and twisted *cro1Δ* cells. White arrowheads mark matured and developing oral structures. Scale bar, 10 μm. **(G)** Twisted cell phenotype is rescued at reduced ciliary forces. To reduce ciliary forces, *cro1Δ* cells were enriched for the twisted cell phenotype at 39°C for 24 h before they were temperature shifted to 25°C. Percentage of twisted cells was assessed 24 h after temperature shift. **(H)** Schematic analysis to quantify SF length and local SF angle ( $\theta$ ) relative to the anterior BB. **(I)** Local SF angle is wider for BBs that exhibit local disorientation as compared to BBs within twisted rows at the elevated force-induced state. **(J)** Locally disoriented BBs possess shorter SFs than BBs in twisted rows.  $n \geq 285$  SFs ( $\geq 37$  cells). **(K)** BB spacing is marginally increased in *cro1Δ* cells at steady-state and elevated force-induced state.  $n \geq 120$  ( $\geq 30$  cells). **(L)** Frequency of SF-pcMT contacts in WT, *cro1Δ*, and *disA-1* cells. Schematic illustrates the position and orientation of two BBs within a region of interest (white boxes). White arrowhead marks BB clusters in *disA-1* mutants.  $n = 240$  SFs (24 cells). Scale bar, 2 μm. tMT, transverse microtubule. Mann-Whitney test. \* denotes  $P < 0.01$ . Mean  $\pm$  SD.

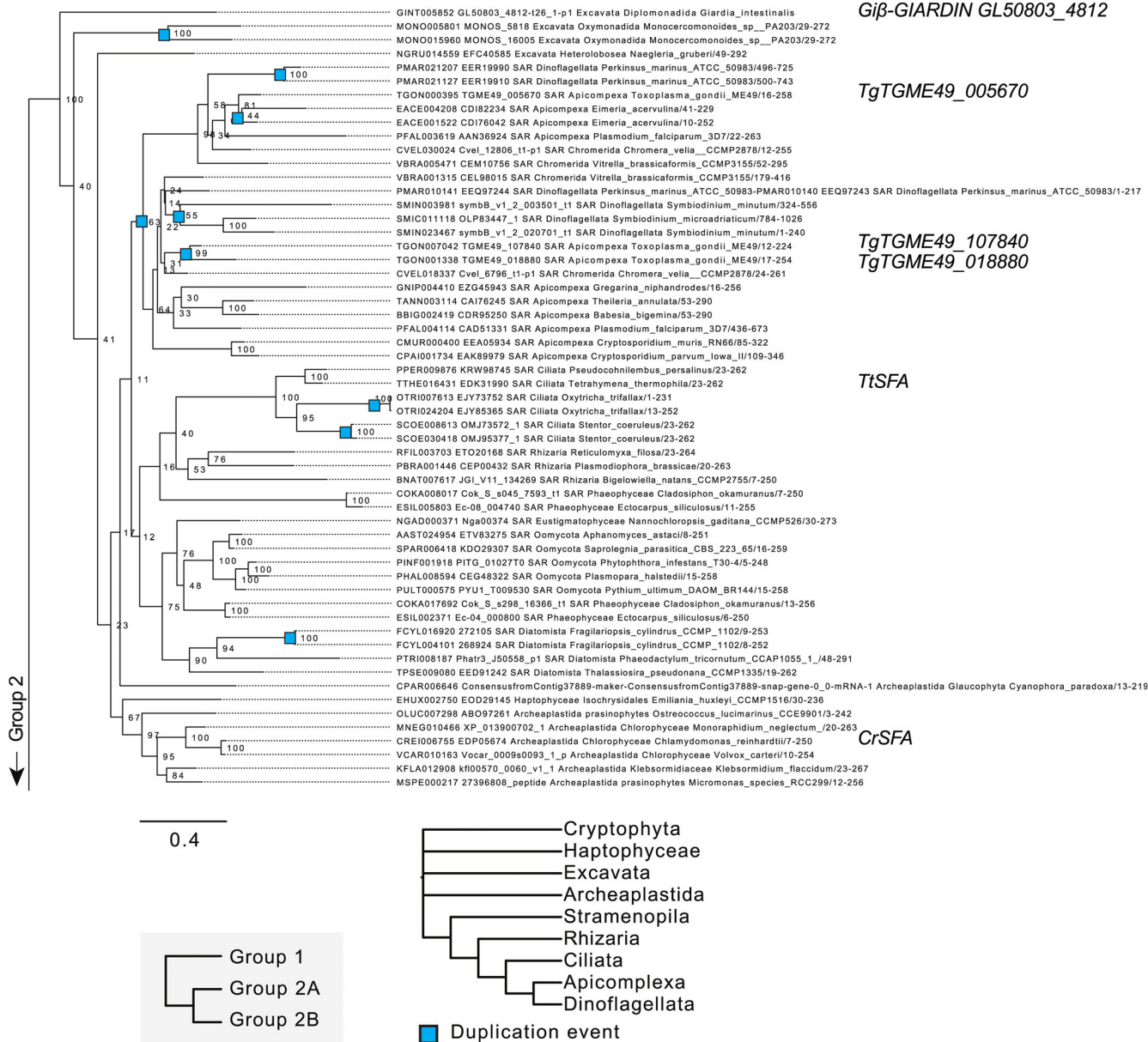
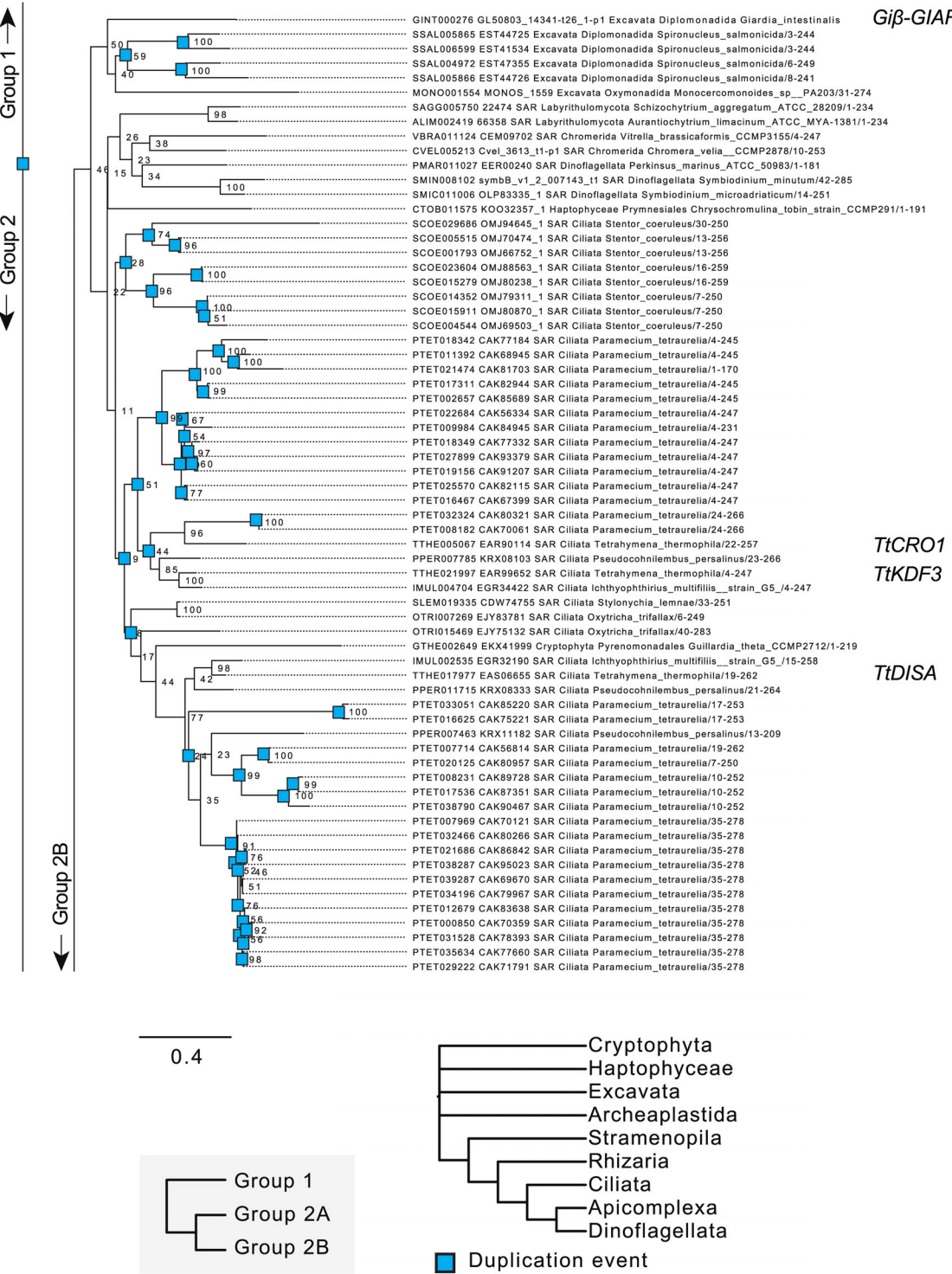
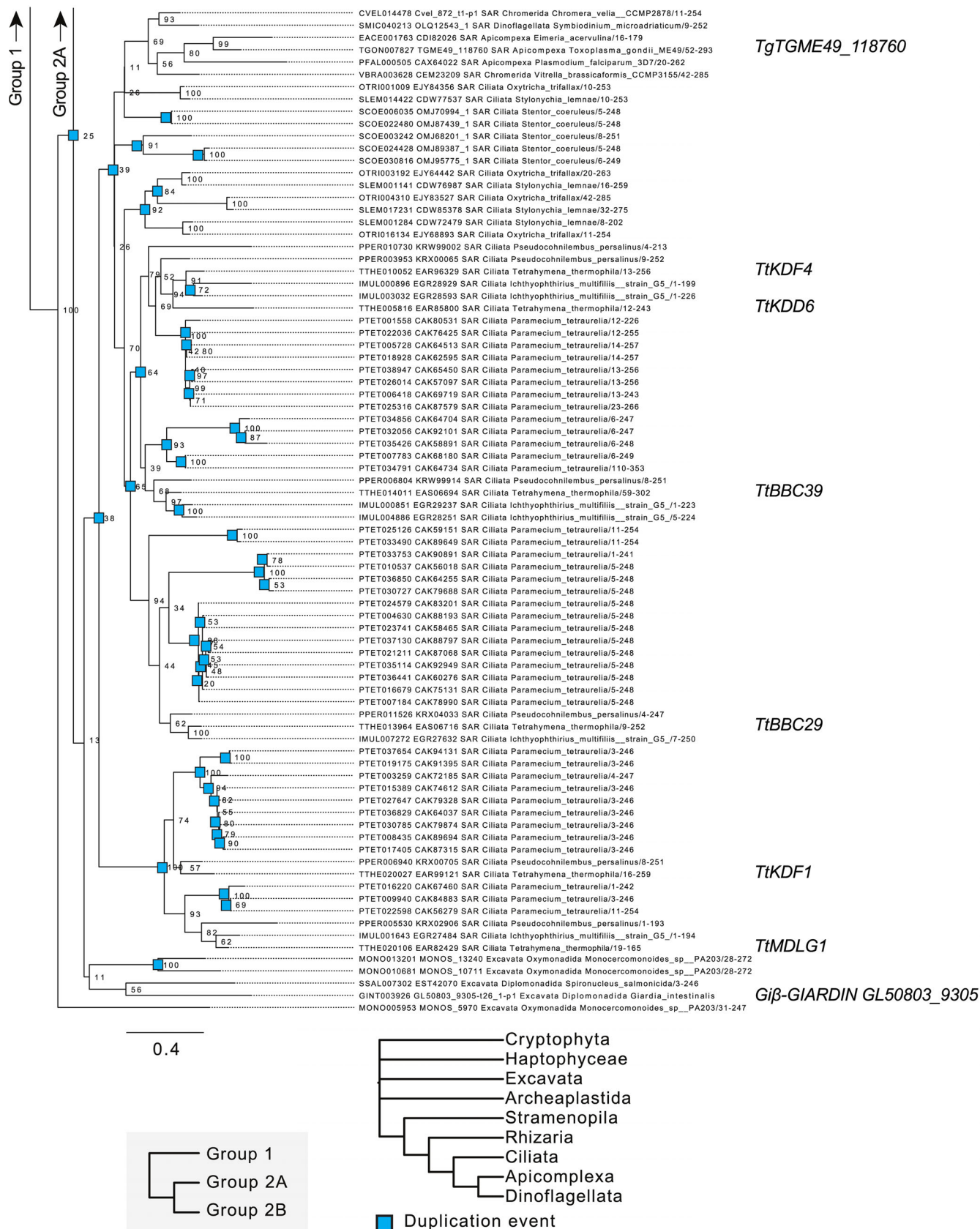


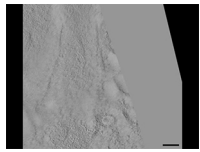
Figure S4. **Phylogenetic analysis of Group 1 SFA homologues among ciliates and algae.** Rooted phylogenetic tree of 205 SFA protein sequences gathered from 50 species of protists and algae. Figure only indicates SFA proteins that belong to the Group 1 clade.



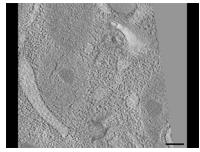
**Figure S5. Phylogenetic analysis of Group 2A SFA homologues among ciliates and algae.** Rooted phylogenetic tree of 205 SFA protein sequences gathered from 50 species of protists and algae. Figure only indicates SFA proteins that belong to the Group 2A clade.



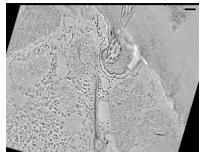
**Figure S6. Phylogenetic analysis of Group 2B SFA homologues among ciliates and algae.** Rooted phylogenetic tree of 205 SFA protein sequences gathered from 50 species of protists and algae. Figure only indicates SFA proteins that belong to the Group 2B clade.



Video 1. **Serial, tomographic slices and model depicting cross section of SF–pcMT interface at steady-state (25°C).** Electron densities link SFs to pcMT bundles. BB, red; SF, green; pcMT, cyan; linkages, purple. Scale bar, 200 nm.



Video 2. **Serial, tomographic slices and model depicting cross section of SF–epiplasm interface at steady-state (25°C).** BB, red; SF, green; epiplasm, white; linkages are absent. Scale bar, 200 nm.



Video 3. **Serial, tomographic slices and model depicting cross section of SF–epiplasm interface at elevated force-induced state (38°C).** Electron densities link SF to the epiplasm. BB, red; SF, green; epiplasm, white; linkages, purple. Scale bar, 200 nm.

**Table S1 shows tabulation of oligo sequences used for fluorescent protein tagging and the functional study of *Tetrahymena* Cro1p. Supplemental data file 1 provides multiple sequence alignment of 205 SFA protein sequences from 50 protist and algae species.**