

1 RESEARCH ARTICLE

2 **Kinesin-5/Cut7 moves bidirectionally on fission-yeast spindles
3 with activity that increases in anaphase**

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6

7 **ABSTRACT**

8 Kinesin-5 motors are essential to separate mitotic spindle poles and assemble a bipolar spindle in many organisms. These motors
9 crosslink and slide apart antiparallel microtubules via microtubule plus-end-directed motility. However, kinesin-5 localization is
10 enhanced away from antiparallel overlaps. Increasing evidence suggests this localization occurs due to bidirectional motility or
11 trafficking. Purified fission-yeast kinesin-5 Cut7p moves bidirectionally, but bidirectionality has not been shown in cells and the
12 function of the minus-end-directed movement is unknown. We characterized the motility of Cut7p on bipolar and monopolar spindles
13 and observed movement toward both plus and minus ends of microtubules. Notably, the activity of the motor increases at anaphase
14 B onset. Perturbations to microtubule dynamics only modestly changed Cut7p movement, while Cut7p mutation reduced movement.
15 These results suggest that the directed motility of Cut7p contributed to the movement of the motor. Comparison of Cut7 mutant
16 and human Eg5 localization suggest a new hypothesis for the function of minus-end-directed motility and spindle-pole localization
17 of kinesin-5s.
18

19 **KEYWORDS:** mitosis, kinesin-5, cut7, fission yeast

20

21 **INTRODUCTION**

22 While kinesin-5 motors have long been known to play an important role in bipolar mitotic spindle assembly and chromosome segregation,
23 the links between kinesin-5 motility, localization, and force generation remain incompletely understood. Kinesin-5 motors are essential
24 for mitotic spindle formation in many organisms because they separate spindle poles to build a bipolar spindle (Enos and Morris (1990);
25 Hagan and Yanagida (1990); Hoyt et al. (1992); Sawin et al. (1992); Blangy et al. (1995), Fig. 1). Kinesin-5s are homo-tetramers, with
26 two dimeric motors linked antiparallel by a central minifilament (Cole et al. (1994); Kashina et al. (1996); Gordon and Roof (1999);
27 Acar et al. (2013); Scholey et al. (2014); Singh et al. (2018), Fig. 1A,B). Thus kinesin-5 can crosslink antiparallel microtubules (MTs)
28 and slide them apart, both in cells and in reconstituted systems (Sharp et al., 1999; Kapitein et al., 2005; Hildebrandt et al., 2006; Tao
29 et al., 2006; van den Wildenberg et al., 2008; Shimamoto et al., 2015; Bodrug et al., 2020). This activity contributes both to spindle pole
30 separation as the bipolar spindle forms and to spindle elongation in anaphase B (Goshima and Scholey, 2010; Mann and Wadsworth,
31 2018; Scholey et al., 2016). Antiparallel sliding is therefore crucial to kinesin-5 function in mitosis, and depends on the motor's motility
32 toward MT plus ends (Fig. 1C). Consistent with this view, kinesin-5 depletion or inhibition leads to monopolar spindles (Hagan and
33 Yanagida, 1990; Sawin et al., 1992; Mayer et al., 1999).

34 However, more recent results have called into question whether the elegantly simple antiparallel-sliding model can fully explain
35 kinesin-5 function. Antiparallel sliding requires antiparallel MTs, which are most abundant near the center of the spindle, where micro-
36 tubules from both poles interdigitate (McIntosh and Landis, 1971; Ding et al., 1993). However, kinesin-5s localize more strongly near
37 spindle poles in many organisms (Fig. 1), including *Schizosaccharomyces pombe* (Hagan and Yanagida, 1992), *Saccharomyces cere-
38 visiae* (Shapira et al., 2017), *Xenopus laevis* (Sawin et al., 1992; Cahu et al., 2008), and mammalian (Blangy et al., 1995; Gable et al.,
39 2012) cells. We have limited understanding of how pole localization occurs for a plus end-directed motor, but there are three broad ideas
40 for the mechanism: trafficking by a minus-end-directed motor that carries kinesin-5 as cargo, intrinsic bidirectional motility of kinesin-5
41 that moves motors toward spindle poles, and direct pole binding.

42 Minus-end-directed trafficking occurs for vertebrate kinesin-5/Eg5, which is plus-end directed but can be transported toward MT
43 minus ends by dynein (Uteng et al., 2008; Gable et al., 2012). The trafficking is regulated by TPX2 (Ma et al., 2010; 2011; Gable et al.,
44 2012; Balchand et al., 2015). Importantly, chimeric kinesin-5s made from the Eg5 tail and the motor of other kinesins fail to assemble
45 a bipolar spindle, even though these motors localize to spindle MTs (Cahu and Surrey, 2009). This evidence suggests that bidirectional
46 trafficking of Eg5 by dynein may play a role in spindle assembly.

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47 Both budding- and fission-yeast kinesin-5s have intrinsic bidirectional motility on MTs *in vitro* (Roostalu et al., 2011; Gerson-Gurwitz
48 et al., 2011; Avunie-Masala et al., 2011; Thiede et al., 2012; Fridman et al., 2013; Edamatsu, 2014; 2016; Britto et al., 2016). For
49 budding-yeast kinesin-5/Cin8, the average direction of motion changes with solution conditions and mutations (Gerson-Gurwitz et al.,
50 Roostalu et al., 2011; Shapira and Gheber, 2016; Shapira et al., 2017). Other work suggests that crowding of the MT lattice and
51 motor clustering affect kinesin-5 speed and directionality (Britto et al., 2016; Shapira et al., 2017; Bodrug et al., 2020).

52 The kinesin-5 C-terminal tail contributes to spindle and spindle-pole localization. The tail contains conserved phosphorylation sites
53 (Heck et al., 1993; Sawin and Mitchison, 1995; Blangy et al., 1995; Drummond and Hagan, 1998; Cahu et al., 2008; Rapley et al., 2008;
54 Akera et al., 2015). Truncation or mutation of the tail can decrease or eliminate spindle (Sawin and Mitchison, 1995) or pole (Sharp
55 et al., 1999; Olmsted et al., 2014) localization. Further, tail truncation in the metazoan kinesin-5 Eg5 significantly impairs microtubule
56 crosslinking and sliding (Weinger et al., 2011; Bodrug et al., 2020), leading to greatly reduced sliding force generation (Bodrug et al.,
57 2020). In budding yeast, deletion of the kinesin-5/Cin8 tail is lethal if the second kinesin-5/Kip1 is absent (Hildebrandt et al., 2006),
58 consistent with idea that tail deletion impairs the ability of Cin8 to separate spindle poles. In fission yeast, kinesin-5/Cut7 pole localization
59 occurs at least in part by direct binding of the tail to γ -tubulin at spindle poles (Olmsted et al., 2014). Recent work has suggested a
60 structural mechanism by which tail-motor interactions could slow ATP hydrolysis (Bodrug et al., 2020). However, the mechanisms by
61 which the C-terminal tail and its phosphorylation affect kinesin-5 motors are incompletely understood.

62 Previous computational modeling of *S. pombe* spindle assembly suggested that kinesin-5/Cut7 bidirectionality may localize it at
63 spindle poles for proper spindle assembly (Blackwell et al., 2017; Edelmaier et al., 2020). Simulated bipolar spindles only assembled
64 when kinesin-5 moved bidirectionally. The minus-end directed movement enhanced the motors' spindle-pole localization in the model,
65 positioning them to generate force and separate spindle poles (Blackwell et al., 2017; Edelmaier et al., 2020). Experimental work on
66 budding-yeast kinesin-5 reached a similar conclusion (Shapira et al., 2017). However, direct experimental test of these model predictions
67 in fission yeast has been limited because kinesin-5/Cut7 bidirectional motility has previously been observed only for the purified motor
68 (Edamatsu, 2014; 2016; Britto et al., 2016). Furthermore, we currently do not understand why (or whether) bidirectional motility and
69 pole localization are beneficial in mitosis, and whether bidirectional trafficking/motility of kinesin-5 may be altered at different times in
70 mitosis.

71 Cut7p in the fission yeast *Schizosaccharomyces pombe* is a useful model for an in-depth study of this bidirectional kinesin on the
72 spindle. This kinesin-5 is well-established as intrinsically bidirectional *in vitro* (Edamatsu, 2014; 2016; Britto et al., 2016). Cut7p is the
73 sole kinesin-5 motor in this organism. It is essential in most genetic backgrounds (Hagan and Yanagida, 1990), but as in other cells, its
74 loss can be rescued by deletion of this cell's two kinesin-14s (Pidoux et al., 1996; Troxell et al., 2001; Rincon et al., 2017; Yukawa et al.,
75 2018; Lamson et al., 2020; Yukawa et al., 2020). Moreover, Cut7p shares many properties with kinesin-5s more generally, including its
76 basic domain structure and localization (Fig. 1). Study of directional motility on the spindle can take advantage of spindle-pole body
77 (SPB) insertion defects in *cut11-ts* cells (West et al., 1998), which lead to monopolar spindles in cells that contain only one active SPB.
78 In mitosis of these cells at restrictive temperature, a spindle MT bundle forms in which all the plus-ends are distal to the single pole,
79 allowing facile assignment of motor movement to polarity (Akera et al., 2015). Fission yeast spindle microtubules appear not to undergo
80 poleward flux. This has been demonstrated by photobleaching of anaphase spindles (Mallavarapu et al., 1999; Sagolla et al., 2003),
81 and likely is true throughout mitosis due to the capping of spindle microtubule minus-ends with γ -tubulin. The lack of poleward flux
82 simplifies the analysis of motor movement on the spindle. Further, *S. pombe* monopolar spindles can enter anaphase with spindle MT
83 elongation occurring at similar speeds as in bipolar spindles (Masuda et al., 1992; Krüger et al., 2021), facilitating study of changes in
84 kinesin-5 motility before and after anaphase onset. Fission yeast therefore allow study of significant aspects of kinesin-5 function in a
85 cell type that is suitable for detailed analysis.

86 RESULTS

87 Cut7p localizes to mitotic spindle microtubules and poles in *cut11+* and *cut11-7* cells

88 To study kinesin-5/Cut7 localization and motility on fission-yeast spindles, we constructed strains containing Cut7-GFP and low-level
89 mCherry-tubulin to label spindle microtubules (Snaith et al. (2010); Blackwell et al. (2016), Fig. 1D-F, Methods). In some cells, we used
90 SPB component Sid4-mCherry to mark spindle poles (Chang and Gould, 2000). *S. pombe* *cut7* fused to 1 or 3 GFP molecules at its
91 C-terminal tail can replace the endogenous motor (Fu et al., 2009), and we used both constructs in this study. For simplicity in the text
92 both constructs are referred to as Cut7-GFP. The exact label is noted in the figure panels and legends. The GFP-tagged motor is functional
93 for spindle assembly with no observed growth or mitotic defects, and its localization is similar to the endogenous untagged motor (Hagan
94 and Yanagida, 1992). This is consistent with observations of other GFP-tagged kinesin-5 motors (Avunie-Masala et al., 2011; Shapira
95 et al., 2017). Consistent with previous studies (Hagan and Yanagida, 1992; Yukawa et al., 2015), we observed Cut7-GFP localized to
96 the unseparated spindle poles early in mitosis, and also to the interpolar spindle as mitosis progressed (Fig. 1D-F). We observed similar
97 behavior in *cut11+* cells (Fig. 1E), and in temperature-sensitive *cut11-7* cells (West et al., 1998; Akera et al., 2015) grown at permissive
98 temperature (Fig. 1F).

99 Cut7p moves bidirectionally on bipolar spindles with activity that increases in anaphase B

100 To better assess Cut7p localization and motility during mitosis, we constructed kymographs of Cut7-GFP on bipolar spindles (Fig. 1G,H,
101 2). Two-color kymographs with Sid4-mCherry labeling SPBs (Fig. 1G) or mCherry-atb2 labeling spindle microtubules (Fig. 1H) both
102 showed localization of Cut7-GFP near spindle poles throughout mitosis. Therefore, we constructed additional kymographs for analysis
103 using the GFP signal only, where bright spots of cut7-GFP labeled the ends of the spindle (Fig. 2A-H). As the spindle poles separated,
104 bright clusters of Cut7-GFP appeared on the interpolar spindle. Diagonal streaks of Cut7-GFP signal were visible in the kymographs,

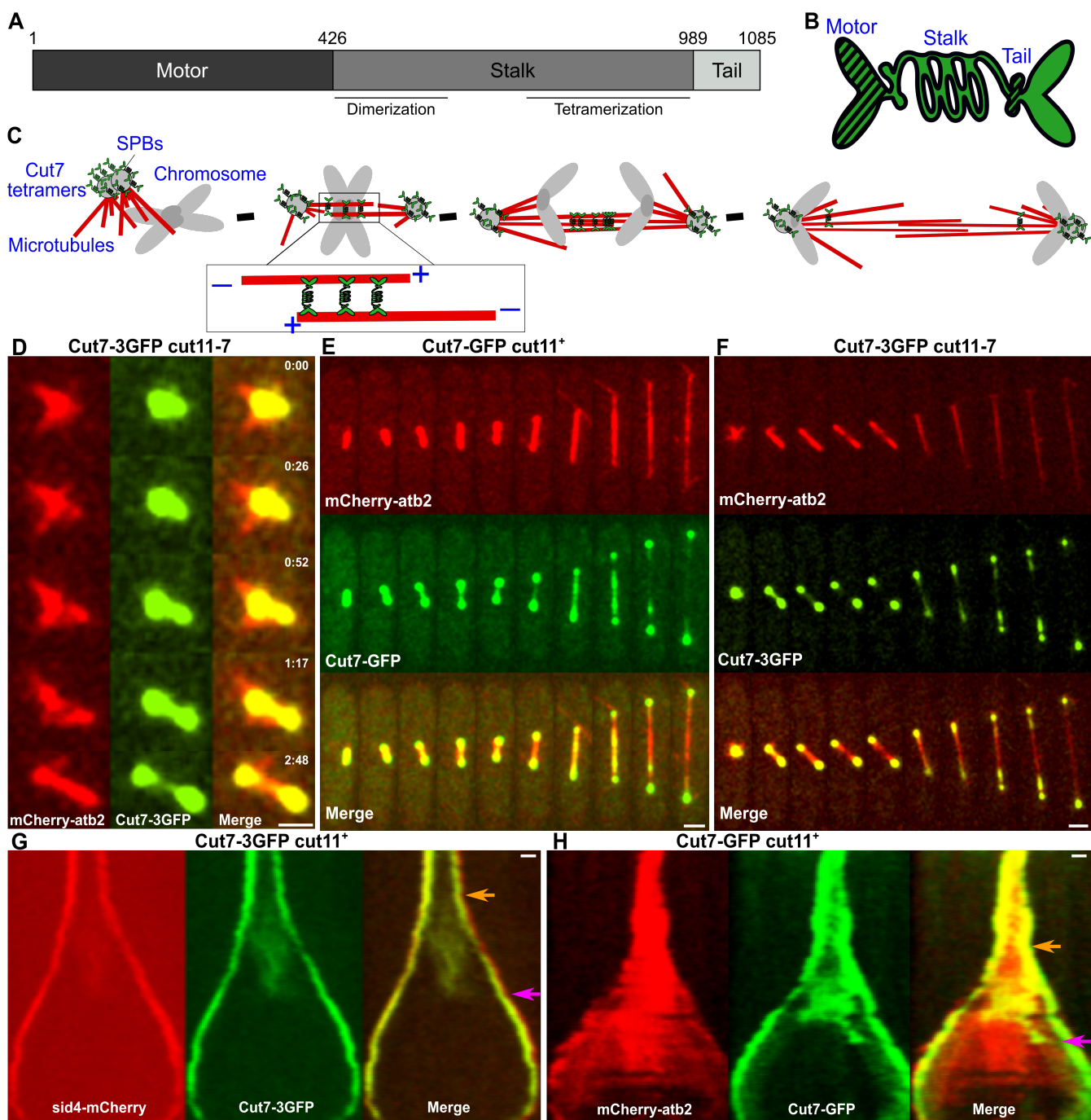


Fig. 1. Overview of Cut7p structure and localization (A) The major domains of Cut7p delimited by amino acid location. (B) Cartoon of a Cut7p tetramer, showing its 2-fold symmetry. (C) Schematic of localization of Cut7p on a mitotic spindle in *S. pombe*. Inset shows Cut7p crosslinking antiparallel MTs. (D) Cut7-3GFP spindle localization in a temperature-sensitive *cut11-7* strain at permissive temperature. As the monopolar spindle becomes bipolar, a subset of Cut7-3GFP (green) moves from the MTs (red) near the left SPB to the MTs near the right SPB. Scale bar: 1 μ m. (E-F) Example of *cut7-GFP*, *cut11⁺* (E) and *cut7-3GFP*, *cut11-7* (F) cells showing Cut7-GFP (green) localization on MTs (red) throughout mitosis. Scale bars: 2 μ m. (G) Two-color kymograph of Sid4-mCherry (left), Cut7-3GFP (center), and merge (right). (H) Two-color kymograph of mCherry-atb2 (left), Cut7-GFP (center), and merge (right). Orange arrows mark anaphase B onset; magenta arrows mark the time when visible Cut7-3GFP disappears from the interpolar spindle. Scale bars: 1 μ m. All images taken at 25 °C.

which represent directional movement of one or more Cut7 motors (Fig. 2A-H, S2). These directed movements were poleward (toward the nearest spindle pole, green arrowheads) and antipoleward (away from the nearest spindle pole, red arrowheads) on bipolar spindles (Fig. 2A-H). To more easily visualize individual or small clusters of Cut7-GFP, we used both photobleaching and photoactivation. After activation of a subset of photoactivatable Cut7-PA-GFP (marked by blue arrowhead on left SPB in figure 2E), we were able to observe

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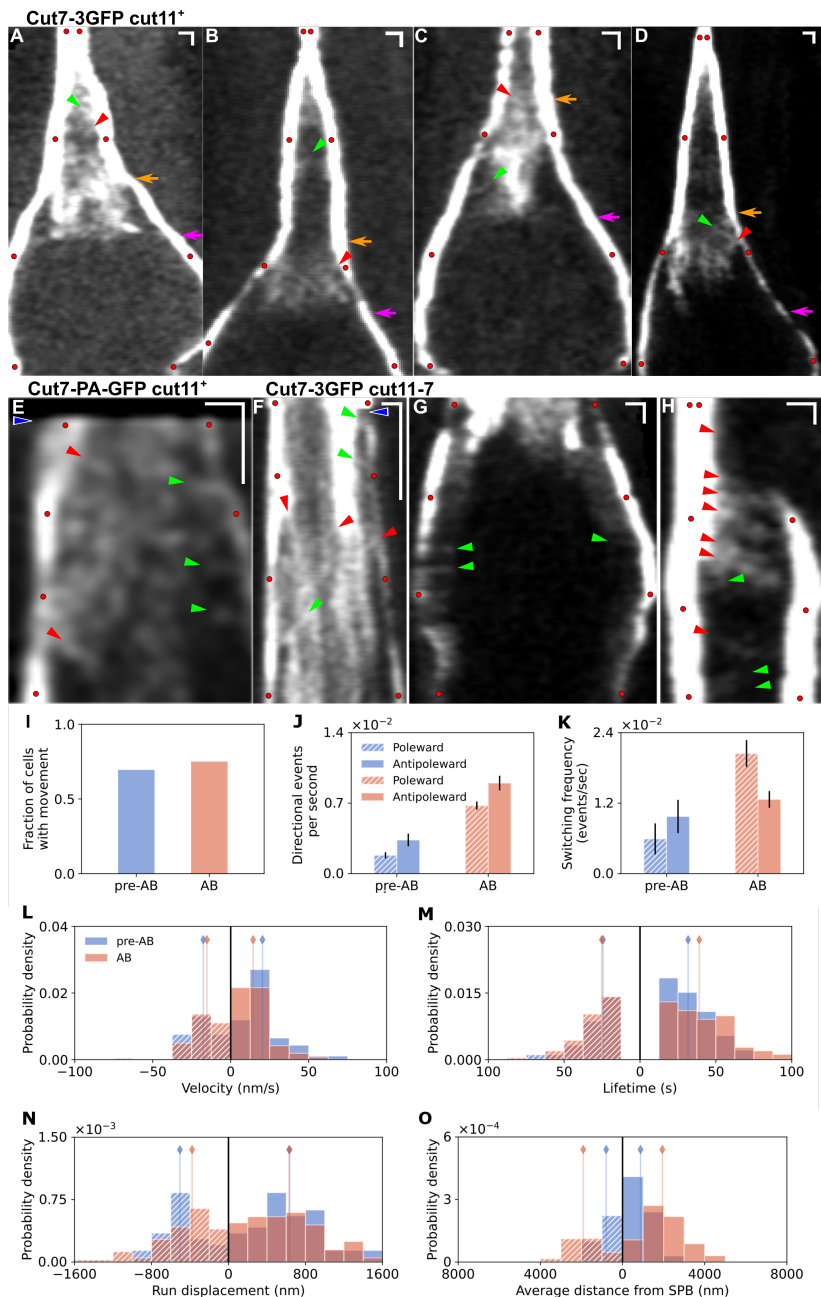


Fig. 2. Localization and motility of Cut7p on bipolar spindles. (A-D) Kymographs of Cut7-3GFP on bipolar spindles from the onset of mitosis through the end of anaphase. Time reads from top to bottom. Orange arrowheads mark anaphase B onset; magenta arrowheads mark the time when visible Cut7-3GFP disappears from the interpolar spindle. Red arrows mark anti-poleward movement and green arrowheads mark poleward movement (relative to the nearest SPB). (E) Photoactivated-GFP kymograph of Cut7-GFP during anaphase B. The blue arrowhead marks the time of photoactivation near the left SPB. (F) Kymograph with fluorescence photobleaching of Cut7-GFP during anaphase B. The blue arrowhead marks the photobleaching event near the right spindle pole. Bright tracks (likely clusters of Cut7-GFP) and dim tracks (likely small clusters or individual motors) are visible. Near the photobleached pole, both poleward and anti-poleward movement is visible. (G) Kymograph in late anaphase B showing poleward movement of Cut7-3GFP that remains on the spindle. (H) Cut7-3GFP localization during spindle pole separation at the onset of mitosis. Cut7-3GFP moves from the left spindle pole and accumulates on the right spindle pole, though the final brightness of the right pole is great enough to suggest that Cut7-3GFP also likely binds directly from the nucleoplasm. SPBs: red circles; Vertical scale bars: 60 s; Horizontal scale bars: 1 μ m. (I–K) Comparison of Cut7-GFP motility before and after the onset of anaphase B. For quantification, see tables S1 and S2. (I) Fraction of cells in which visible movement occurred. (J) Frequency of visible directed movement. (K) Switching frequency out of each directed event. (L–O) Quantification of Cut7-GFP motility events. Values corresponding to poleward-directed movement are on the left side of the axis, anti-poleward directed movement on the right. Vertical bars represent median of the distribution. For quantification, see tables S1 and S2. (L) Velocity. (M) Lifetime. (N) Run displacement. (O) Average distance from the spindle pole including all points from each event.

109 poleward and anti-poleward movement of Cut7-GFP. Photobleaching near one spindle pole (marked by blue arrowhead near right SPB
110 in fig. 2F) allowed additional observation of poleward and anti-poleward movement of Cut7-GFP near that SPB.

We identified the onset of anaphase B as the time when the spindle elongation rate began to increase (fig. 1G,H, orange arrows, fig. 2A-D, Krüger et al. (2021)). Typically the intensity of Cut7-GFP fluorescence near the spindle midzone was highest just after anaphase B onset. In early anaphase B, the kymographs suggested that Cut7p binding and/or movement increased, because a relatively bright, diffuse region of Cut7-GFP intensity appeared (fig. 1G,H, fig. 2A-D, between orange and magenta arrows). Subsequently, most Cut7-GFP intensity disappeared from the interpolar spindle (fig. 1G,H, fig. 2A-D, magenta arrows). After this time, any Cut7-GFP signal visible in the interpolar spindle typically moved toward the nearest spindle pole (Fig. 2G). The midzone of mid-to-late anaphase spindles contained little visible Cut7-GFP, consistent with previous findings that kinesin-6/Klp9 contributes more than Cut7p to anaphase B spindle elongation in fission yeast (Yukawa et al., 2018; 2019).

The *S. pombe* interpolar spindle in early mitosis includes some MTs that extend from one pole all the way to the other (Ding et al., 1993; Ward et al., 2015). As a result, poleward and anti-poleward movement cannot be assigned to plus- or minus-end-directed movement. However, in late anaphase, MTs visible in spindle electron micrographs usually overlap only near the spindle midzone (Ding et al., 1993; Ward et al., 2015). Therefore, the poleward Cut7-GFP movement visible near the spindle poles in late anaphase (Fig. 2G) could be minus-end-directed. This suggests that Cut7p can move toward the minus-ends of spindle MTs, consistent with its activity *in vitro* (Edamatsu, 2014; 2016; Britto et al., 2016).

We observed that Cut7-GFP was typically visible near one spindle pole earlier than the other. In *cut11-7* cells that have SPB insertion defects at restrictive temperature, we observed that even at permissive temperature, the asymmetry in early Cut7-GFP localization was exaggerated (Fig. 2H). Perhaps this is due to a delay in MT nucleation at the second SPB as observed previously for *cut11-6* (Zhang and Olierenko, 2014). As a result, we observed bright Cut7-GFP signal near one spindle pole and little signal on the other, shortly after the poles separated (Fig. 2H). The kymograph shows that Cut7-GFP traveled primarily toward the dimmer spindle pole. In addition, the signal at the dimmer pole increases rapidly (Fig. 2H). This suggests that some Cut7-GFP pole accumulation likely occurs due to direct binding of the motor to spindle poles, consistent with previous work (Olmsted et al., 2014). In addition, these results suggest that Cut7-GFP moves in both directions between the spindle poles as the bipolar spindle assembles, and that Cut7-GFP signal can increase near a spindle pole due to both movement from the interpolar spindle and direct binding.

To further analyze Cut7p movement, we created custom Matlab software for kymograph analysis (Methods). We hand-traced visible Cut7-GFP tracks and then analyzed events (segments of tracks) that exhibited movement poleward or anti-poleward, or pauses (Fig. S1, S2). Additionally, we divided events into pre- and post-onset of anaphase B. For each movement event, we analyzed the velocity, lifetime, run displacement, and distance from the closest spindle pole. From the entire data set we identified the total number of cells that showed identifiable directed Cut7-GFP tracks, the frequency at which directional events occurred, and the state exit frequency (the switching frequency, defined as the rate at which a directional event ends by changing direction or pausing, Table S1, S2).

Consistent with visual inspection of the kymographs, Cut7p activity significantly increased after the onset of anaphase B (Fig. 2I-K). Before and after anaphase onset, the overall fraction of cells with bipolar spindles imaged that showed visible Cut7-GFP movement was similar, 69% pre-anaphase and 75% in anaphase B (Fig. 2I, Table S1). Despite this similarity, the number of directional events observed per second was larger by a factor of 2-5 after anaphase onset (Fig. 2J). Both pre-anaphase B and in anaphase B, anti-poleward events occurred slightly more frequently than poleward, consistent with the general tendency of Cut7-GFP to accumulate near spindle poles (Fig. 2J). Next, we measured the rate at which directed events ended (switching frequency), either due to pausing or changing direction (Fig. 2K). This rate increased slightly for anti-poleward events but by a factor of 3 for poleward events in anaphase B, meaning that direction switching and pausing occurred more frequently. Prior to anaphase B, poleward movement had a slightly lower state exit frequency (Fig. 2K), meaning that poleward movement persisted for longer than anti-poleward. During anaphase B, the state exit frequency of poleward movement was almost twice that of anti-poleward, meaning that anti-poleward movement lasted longer (was more persistent). Therefore, during early anaphase Cut7p movement events both occurred more frequently and switched direction or paused more frequently.

Analysis of individual directed movement events allowed us to determine motility parameters of poleward and anti-poleward events, both pre- and during anaphase B (Fig. 2L-O, Table S2). Note that to facilitate comparison of poleward and anti-poleward movement, the left half of the graph/negative numbers show data for poleward movement, while the right half of the graph/positive numbers denote anti-poleward movement (always determined relative to the closest pole at the start of the event). The median speed of Cut7-GFP movement was 15-20 nm s⁻¹ for both directions of movement and stages of mitosis (Fig. 2L). Therefore the change in Cut7p activity in anaphase does not appear to be driven by a change in motor speed. The lifetime of directed events tended to be longer for anti-poleward movement (Fig. 2M). The run displacement was lower for poleward movement and decreased in anaphase B, while for anti-poleward events it was higher and stayed the same in anaphase B (Fig. 2N). By averaging all points from all tracks, we measured the average distance of visible tracks from the closest spindle pole and found that this increases by a factor of two in anaphase, reflecting a greater likelihood of Cut7 tracks to be visible farther from the spindle poles in anaphase B (Fig. 2O). We found no significant correlation between Cut7-GFP spot intensity and particle velocity, lifetime, or run length (Fig. S3, pink).

In summary, Cut7-GFP moved in both directions along bipolar spindles, and its direction-switching activity increased significantly in anaphase B. The speed of directed events was typically 15-20 nm s⁻¹ and lifetime 10-15 sec. Our observation of Cut7-GFP moving both poleward and antipoleward on bipolar spindles and switching direction was consistent with bidirectional motility on spindle MTs. An alternative way to interpret these movements is to attribute them to Cut7-GFP remaining at plus ends of growing and shrinking microtubules. Therefore we compared the Cut7-GFP event speeds we measured to MT dynamics measurements. Recent work quantified MT dynamics in fission-yeast bipolar spindles in anaphase and found a growth speed of 23 nm s⁻¹ and shrinkage speed of 68 nm s⁻¹ (Lera-Ramirez et al., 2022). The shrinkage speed is significantly faster than Cut7-GFP poleward movement events that we measured, suggesting that these events likely do not result from motors tracking the plus-ends of depolymerizing MTs. However, the microtubule growth speed is comparable to the Cut7-GFP movement speed we measured. Therefore, the possibility that Cut7 tracks plus-ends of

172 growing MTs is consistent with our measurements. A potentially complicating factor in assessing directionality of movement is that
173 interpolar spindle MTs are mixed in polarity, making it impossible to assign observed events unambiguously as directed toward plus or
174 minus ends of MTs in the diffraction-limited spindle. However, in late anaphase B, results from both electron and light microscopy show
175 that MTs typically end near the midzone: it is rare for a MT from one spindle pole to extend nearly the length of the spindle to end near
176 the opposite spindle pole (Ding et al., 1993; Ward et al., 2015; Lera-Ramirez et al., 2022). Therefore, it is likely (but not guaranteed)
177 that in late anaphase motor movement near the poles is plus-end-directed when anti-poleward and minus-end directed when poleward.
178 To more definitively correlate Cut7p movement with MT polarity, we built on the observation that *S. pombe* monopolar spindles have
179 minus-ends at the SPBs and plus-ends pointing outward. Therefore, we sought to examine Cut7-GFP on monopolar mitotic spindles
180 to determine whether the motor moves both toward plus and minus ends of microtubules.

181 **Cut7p moves bidirectionally on *cut11-7* monopolar spindles**

182 Monopolar mitotic spindles with only one active SPB are observed reproducibly in fission yeast with temperature-sensitive mutations
183 of *cut11* (West et al., 1998). At restrictive temperature, a spindle MT bundle forms in which all the plus ends are distal to the single
184 pole (Fig. 3), allowing unambiguous assignment of motor movement toward MT plus or minus ends (Akera et al., 2015). We therefore
185 tracked Cut7-GFP movement on this unipolar microtubule array to determine whether Cut7-GFP can move toward both the plus and
186 minus ends of spindle MTs. For high time resolution of Cut7-GFP movement, we imaged the mCherry-atb2 channel only every 15
187 timepoints (Fig. S4). We examined these images and chose a MT bundle in the monopolar spindle to construct a kymograph and identify
188 directed Cut7-GFP movement events (fig. 3, S4, S5).

189 In *cut11-7* cells, monopolar spindles initially contained short MTs, similar to bipolar spindles in early mitosis (fig. S4, 3A,B). Later
190 some of the bundles elongated by almost a factor of two (fig. S4, 3C-E). Longer monopolar spindles contained one or more MT bundles
191 that were more stable. The change from short to long monopolar spindles has been shown to occur at the transition into an anaphase-
192 B-like state, based on several cell-cycle markers (Krüger et al., 2021), despite the absence of chromosome segregation in these cells.
193 Based on an average of 10 monopolar spindles that showed rapid elongation during our time of observation, we identified 2.4 μm as the
194 length at which this transition typically occurred, allowing us to define a boundary between spindles pre-anaphase and in anaphase B.
195 This allowed us to compare the movement of Cut7-GFP on these two categories of monopolar spindle.

196 On all monopolar spindles Cut7-GFP localized primarily near the spindle pole, as observed previously (fig. S4, Akera et al. (2015)).
197 In this case, a kymograph of Cut7-GFP signal appeared as a straight vertical bar with few movement events along the microtubule bundle
198 used to construct the kymograph (fig. S4, 3A). On the majority of pre-anaphase monopolar spindles, we observed no motion of Cut7-
199 GFP along MT bundles. However, in some cases we observed short, dim events, likely corresponding to single motors or small clusters,
200 characterized by movement a short distance from the SPB and then back (Fig. 3B). In longer anaphase monopolar spindles, brighter
201 clusters were present either paused or moving along the spindle (Fig. 3C-E, S5). In some cases we photobleached the signal near the
202 SPB to allow visualization of smaller clusters (3C, blue arrow). On these longer monopolar spindles, we often observed Cut7-GFP along
203 spindle MTs more distal to the SPBs (Fig. 3C-E, S5). In some cases, Cut7-GFP moved in distinct tracks originating from the SPB (Fig.
204 S6). For other events, the Cut7-GFP signal appeared brightly at one point along the spindle, without a clear track leading from the SPB
205 (Fig. 3C-E, S5). This may occur due to binding of motors directly from the nucleoplasm.

206 We quantified differences in Cut7-GFP movement on pre-anaphase and anaphase B monopolar spindles, and found that movement
207 of Cut7-GFP away from the SPB occurred more frequently in anaphase B (76% of cells) compared with pre-anaphase spindles (31%,
208 fig. 3F, Table S1). Consistent with this, in anaphase B the frequency of observable directed events was a factor of 1.5-2 larger for both
209 poleward and anti-poleward movement (Fig. 3G). As a result, anaphase B spindles showed significantly more Cut7-GFP tracks than cells
210 pre-anaphase B. The switching frequency out of poleward movement events appeared to increase upon entry to anaphase B (Fig. 3H),
211 consistent with the increase in anti-poleward directed movement.

212 The median velocity increased slightly on monopolar spindles compared to bipolar (Fig. 3I, Table S2), and lifetime and run displacement
213 were somewhat shorter as well (Fig. 3J,K). These changes were small but might reflect differences in Cut7p behavior during parallel
214 versus antiparallel MT crosslinking, as has been shown for the *S. cerevisiae* kinesin-5/Cin8 (Gerson-Gurwitz et al., 2011). The average
215 distance of Cut7-GFP from the SPB was $\approx 1 \mu\text{m}$ (Fig. 3L). Together, these data provide evidence that Cut7p motility on the spindle can
216 be toward either the plus or the minus end of MTs, and there is more motor activity in both directions as cells enter anaphase B.

217 Cut7-GFP on monopolar spindles showed events moving in both directions, consistent with the idea that the motor moves bidirectionally.
218 However, as mentioned above, such bidirectional events could also occur if the motors can track dynamic MT ends. The median
219 speed of directed Cut7-GFP tracks that we measured on monopolar (and bipolar) spindles was 15-20 nm s $^{-1}$ (Fig. 2, 3, Table S2, S3).
220 This is slow compared to previous measurements of fission-yeast spindle MT dynamic instability in monopoles, which found MT growth
221 speed of 45-70 nm s $^{-1}$ and MT shrinking speed of 60-110 nm s $^{-1}$ (Kalinina et al. (2012); Blackwell et al. (2017), Table S3), suggesting
222 that the observed Cut7-GFP movement is unlikely to occur solely to microtubule plus-end tracking. However, it is possible that MTs
223 with slower dynamics could be present in the spindle. Therefore, we perturbed both MT dynamics and Cut7p motility and examined the
224 effects on Cut7-GFP motility on monopolar and bipolar spindles.

225 **Cut7p moves bidirectionally on *klp5 Δ* monopolar spindles that contain more stable microtubules**

226 To test whether the observed bidirectional Cut7p movement could be driven by spindle MT dynamics, we imaged Cut7-GFP on *cut11-7*
227 monopolar spindles in cells containing deletions of kinesin-8/Klp5. Because Klp5p contributes to MT depolymerization, *klp5 Δ* cells
228 have microtubules and spindles that are 2-3 times longer (Fig. 4, right vs. left cartoon) and are more stable and bundled (West et al.,
229 2001). In addition it has been shown that *klp5* deletion causes no change in the rates of microtubule growth and shrinkage; rather it

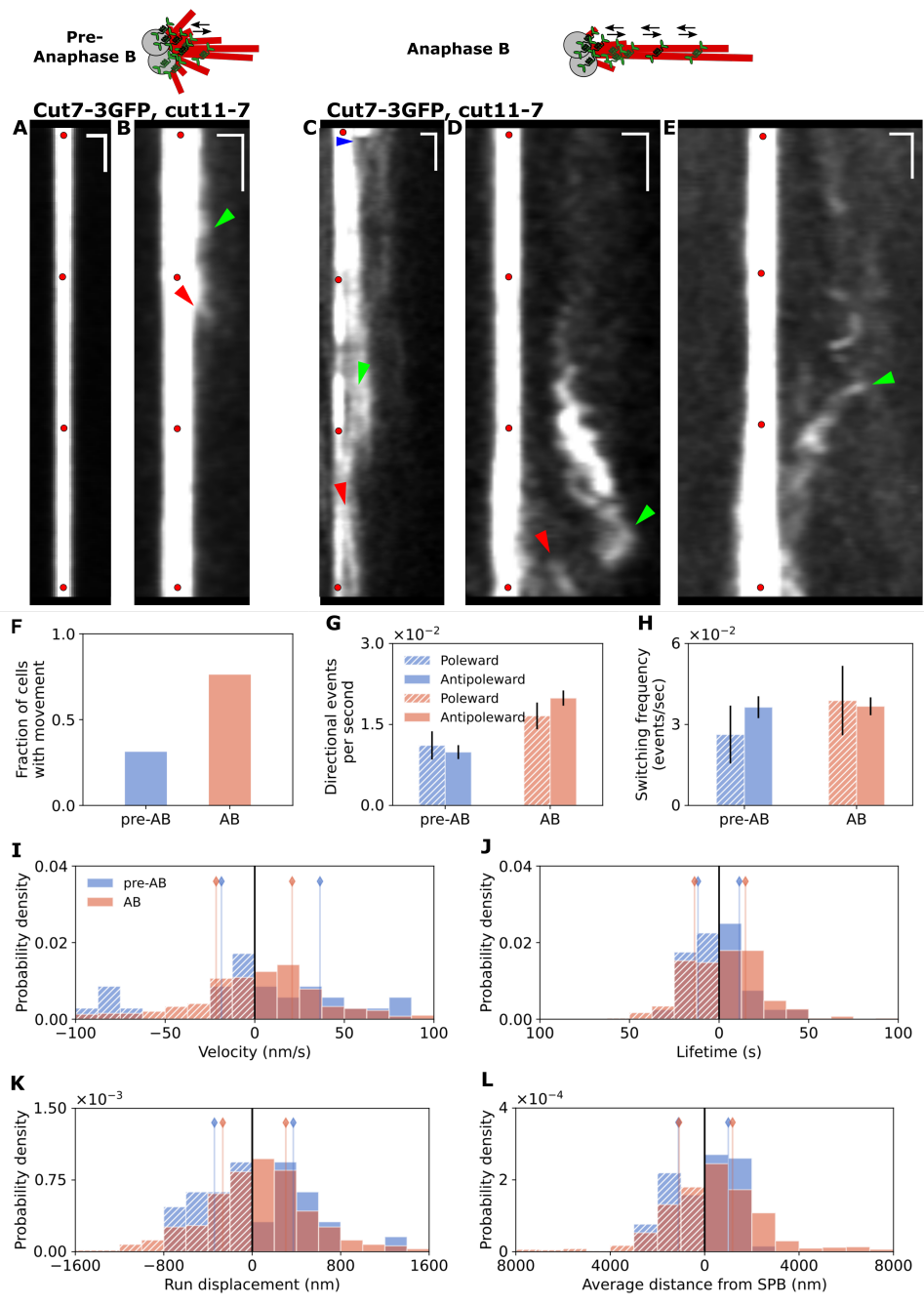


Fig. 3. Localization and motility of Cut7p on monopolar spindles differs in short and long spindles. Schematic of Cut7-GFP movement on monopolar spindles formed in *cut11-7* cells at restrictive temperature. (A-E) Kymographs of Cut7-3GFP on *cut11-7* monopolar spindles. Red arrowheads mark anti-poleward movement and green arrowheads mark poleward movement. (A-B) Pre-anaphase B. (C-E) During anaphase B. (C) Kymograph with photobleaching; the blue arrowhead indicates the time of bleaching the monopolar arm to the right of the spindle pole. SPBs: red circles; Vertical scale bars: 30 s; Horizontal scale bars: 1 μ m. (F-L) Comparison of Cut7-GFP motility before and after the onset of anaphase B. For quantification, see tables S1 and S2. (F) Fraction of cells with visible movement. (G) Frequency of visible directed movement. (H) Switching frequency out of each directed event. (I-L) Quantification of cut7-GFP motility events. Values corresponding to minus-end directed movement are on the left side of the axis, plus-end directed movement on the right. Vertical bars represent median of the distribution. For quantification, see tables S1 and S2. (I) Velocity. (J) Lifetime. (K) Run displacement. (L) Average distance from the SPB including all points from each event.

decreases microtubule dynamicity, by reducing the frequency of both rescue and catastrophe (Unsworth et al., 2008). Consistent with previous work, we observed that monopolar spindles in *klp5 Δ* *cut11-7* cells were significantly longer than those in *klp5 $+$* (Fig. 4A-E, compare Fig. S7 and S4). As in *klp5 $+$* cells most Cut7-GFP on *klp5 Δ* monopolar spindles remained near the SPB (Fig. S7). If Cut7-GFP bidirectional movement occurs primarily due to MT dynamics, we would expect that Cut7p directional movement would occur less frequently and switch direction less frequently in *klp5 Δ* cells compared to *klp5 $+$* cells (Fig. 3).

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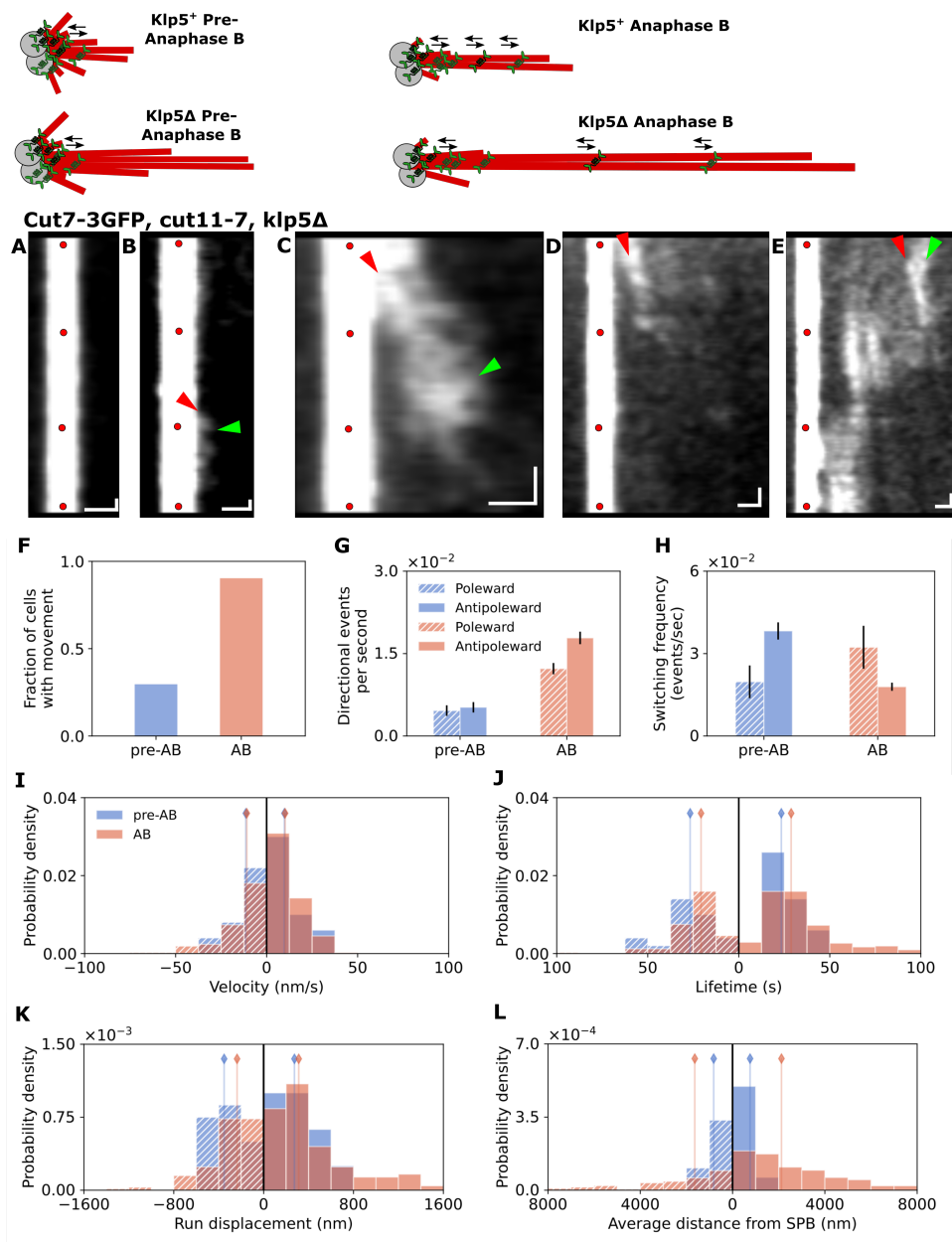


Fig. 4. Longer monopolar spindles make the anaphase B increases in Cut7-GFP motility more evident. Cartoon of Cut7-GFP motor movement on monopolar spindles in *cut11-7* (upper) or in *cut11-7, klp5Δ* cells (lower), before and during anaphase B. (A-E) Kymographs of Cut7-3GFP on *klp5Δ*, *cut11-7* monopolar spindles. Red arrowheads mark anti-poleward movement and green arrowheads mark poleward movement. (A-B) Pre-anaphase B and (C-E) Anaphase B. SPBs: red circles; Vertical scale bars: 30 s; Horizontal scale bars: 1 μ m. (F-L) Comparison of Cut7-GFP motility on *klp5Δ* monopolar spindles before and after the onset of anaphase B. For quantification, see tables S1 and S2. (F) Fraction of cells with any movement. (G) Frequency of visible directed movement. (H) Switching frequency out of each directed event. (I-L) Quantification of cut7-GFP motility events. Values corresponding to minus-end directed movement are on the left side of the axis, plus-end directed movement on the right. Vertical bars represent median of the distribution. For quantification, see tables S1 and S2. (I) Velocity. (J) Lifetime. (K) Run displacement. (L) Average distance from the SPB including all points from each event.

We found that the fraction of pre-anaphase B cells that showed any spindle-associated Cut7-GFP motion was unaffected by the increased spindle length and was still only 30% (Fig. 4F, Table S1, Fig. S8) despite the longer length of spindle MTs on which events could take place. The fraction of cells with observable movement in *klp5Δ* cells increased during anaphase B to 90%. This fraction was higher in *klp5Δ* cells than in *klp5+* cells (76%). This is the opposite of what would be predicted if MT dynamics drive the observed Cut7-GFP movements.

The number of observable directed Cut7-GFP events per second was similar in the *klp5+* and *klp5Δ* monopolar spindles, but with a slight decrease in the pre-anaphase B directed movement event rate (Fig. 4G, Table S1). While the switching rates were comparable in the two strains, the exit rate from anti-poleward movements in anaphase B decreased in *klp5Δ* cells (Fig. 4H, Table S1). This is consistent with the hypothesis that bidirectional movement arises primarily from Cut7-GFP motility. As in *klp5+* cells, motors in pre-anaphase B

klp5 Δ cells switched more frequently to minus-end directed movement, and switched more frequently to plus-end-directed movement in anaphase B (Fig. 4H). Overall, Cut7-GFP movement was not dramatically altered by the changing the length and dynamicity of monopolar spindle MTs (Fig. 4I-L, Table S2). The median velocity of directed events decreased slightly compared to *klp5* $+$, but the velocity maintained a wide distribution (Fig. 4I). The lifetime increased slightly, again with a wide distribution (Fig. 4J). As a result the run length (Fig. 4K) was similar to measurements in *klp5* $+$ monopolar spindles. The average distance from the SPB increased in anaphase spindles, likely because the MTs were longer (Fig. 4L). Together, these data are consistent with the idea that that Cut7p moves bidirectionally on spindle MTs. While movements driven by MT dynamics cannot be completely ruled out, they appear to be a smaller contributor than Cut7p intrinsic motility. To further test whether these movements are due to Cut7p motility, we made several perturbations to Cut7p itself.

Bidirectional motility is altered in Cut7 motor and tail mutants

To test whether perturbation of Cut7p can alter its movement on the spindle, we examined alterations to the motor that affect either the motor domain or the C-terminal tail. First, we asked whether Cut7p with no motor activity would change its localization on the spindle. If Cut7p movements are primarily driven by MT dynamics, we would expect to see a similar pattern of localization in motor-active Cut7p and motor-dead Cut7p. We therefore created a *cut7* mutant with a motor domain that could not bind ATP (*cut7-motor-dead*), constructed by changing three amino acids at residues 164-166. Previous work showed that this mutation blocks Cut7p movement but not microtubule binding (Akera et al., 2015). When *cut7-motor-dead* is the only allele of *cut7* in the cell, it is lethal. Therefore, we expressed it in a strain that also carries the temperature sensitive allele *cut7-446* (Hirano et al., 1986). Of the 25 cells with this genotype we imaged at permissive temperature (25 °C), 2 (8%) formed monopolar spindles and 23 (92%) formed bipolar spindles. Unlike Cut7-GFP (Fig. 1) and Cut7-446-GFP (Fig. 5A), Cut7-motor-dead-GFP was not enriched at spindle poles. Instead, it showed faint diffuse fluorescence along the entire length of the spindle (Fig. 5B). Since Cut7-motor-dead-GFP cannot move along spindle microtubules, the diffuse localization appears to result from direct motor binding both at the spindle midzone and near the poles. To further analyze the localization of Cut7-motor-dead-GFP, we made kymographs of the spindle and motor fluorescence (Fig. S9, Methods). These showed Cut7-motor-dead-GFP spots that colocalized with spindle microtubules, and did not show directed movement along the spindle. Some Cut7-motor-dead-GFP signal was visible near the spindle poles, but did not appear enhanced at the poles as occurs in Cut7-GFP (Fig. 2A-D, S9A,B), or in Cut7-446-GFP (Fig. 5C-F). To compare the localization pattern quantitatively, we selected slices of the kymographs corresponding to specific values of spindle length and averaged GFP intensity across multiple cells with spindles of the same length (Methods). Cut7-GFP showed distinct fluorescence peaks at the ends of the spindle, while Cut7-motor-dead-GFP shows approximately flat intensity distribution along its length (Fig. S9C-F). Therefore, Cut7-motor-dead GFP does not appear to have an increased binding preference for spindle poles. This could occur for at least three reasons: first, the motor-dead mutant could be defective in direct binding to γ -tubulin near the SPBs for an unknown reason. Second, perhaps spindle-pole localization requires higher expression of the motor than occurs for *cut7-motor-dead*. Alternatively, the typical spindle localization of Cut7-GFP may depend at least in part on its minus-end-directed motility. These results, together with our observations of minus-end-directed Cut7-GFP movement toward spindle poles, suggest that motor movement contributes to Cut7-GFP spindle-pole localization. In addition, this result suggests that while MT dynamics may be a contributor to Cut7-GFP movement, Cut7-GFP accumulation at spindle poles does not occur solely due to MT dynamics.

Next, we sought to determine whether the *cut7-446* temperature-sensitive mutation affects Cut7p motility. This is a single point mutation, I954T, near the start of the tail region (Yukawa et al., 2018). As mentioned above, at permissive temperature these cells formed bipolar spindles with Cut7-446-GFP localization similar to Cut7-GFP (Fig. 5A). Further, kymographs of Cut7-446-GFP appeared similar to those of Cut7-GFP throughout mitosis (Fig. 5C-E). We then shifted *cut7-446* cells to restrictive temperature (37 °C), which leads to temperature-sensitive lethality by blocking normal SPB separation (Hagan and Yanagida, 1992; Tallada et al., 2009). In 98% of the monopolar spindles that formed (N=104), Cut7-446-GFP was visible only near the SPBs, with no directed movement events. This suggests that at restrictive temperature the motor either cannot move or only retains minus-end-directed motion; we observed almost no plus-end-directed motion (Fig. 5F), in contrast to *cut11-7* monopolar spindles in which Cut7-GFP showed plus end-directed movements in 36% of spindles (Fig. 3F). This observation is consistent with Cut7p immunofluorescent detection in *cut7-446* cells (Hagan and Yanagida, 1992) and supports that movement of Cut7-GFP is due to its intrinsic motility, and not exclusively by tracking plus-ends of dynamic MTs. In addition, this single amino acid change appeared to abolish or disrupt plus-end movement at restrictive temperature.

To further perturb Cut7p, we truncated the C-terminal tail. The C-terminal tail contains the conserved BimC box containing a Cdk phosphorylation site (Drummond and Hagan, 1998), and point mutations in this region lead to alleles that are temperature sensitive for growth (Rodriguez et al., 2008; Akera et al., 2015). Further, tail truncation in other kinesin-5s can impair MT crosslinking and sliding (Hildebrandt et al., 2006; Bodrug et al., 2020). Therefore we asked whether truncation of the tail at the end of the predicted coiled-coil domain (Fig. S10) affects Cut7p motility. The truncation we constructed removed 96 amino acids from the tail beyond amino acid 988. This mutation appeared to be lethal, as we were unable to produce transformants with *cut7-988* as the sole *cut7* allele in the cell. This is consistent with observations in budding yeast that deletion of the kinesin-5/Cin8 tail is lethal in the absence of kinesin-5/Kip1 (Hildebrandt et al., 2006). To allow observation of motility of Cut7-988-GFP, we examined its phenotype in cells with deletions of *pkl1* and *klp2*, genes that encode two minus-end-directed kinesin-14s. Because Pkl1p and Klp2p are antagonistic to Cut7, deleting them allows study of lethal alleles of *cut7* (Pidoux et al., 1996; Troxell et al., 2001; Rincon et al., 2017; Yukawa et al., 2018; Lamson et al., 2020; Yukawa et al., 2020). This combination of mutations produced viable cells at 25°C, but monopolar spindles at 37°C.

On bipolar spindles, Cut7-988-GFP localized predominantly to the spindle poles, similarly to Cut7-GFP (Fig. 5G-I). This suggests that the motor has sufficient activity to move to spindle poles or has a binding preference near microtubule minus ends, in contrast

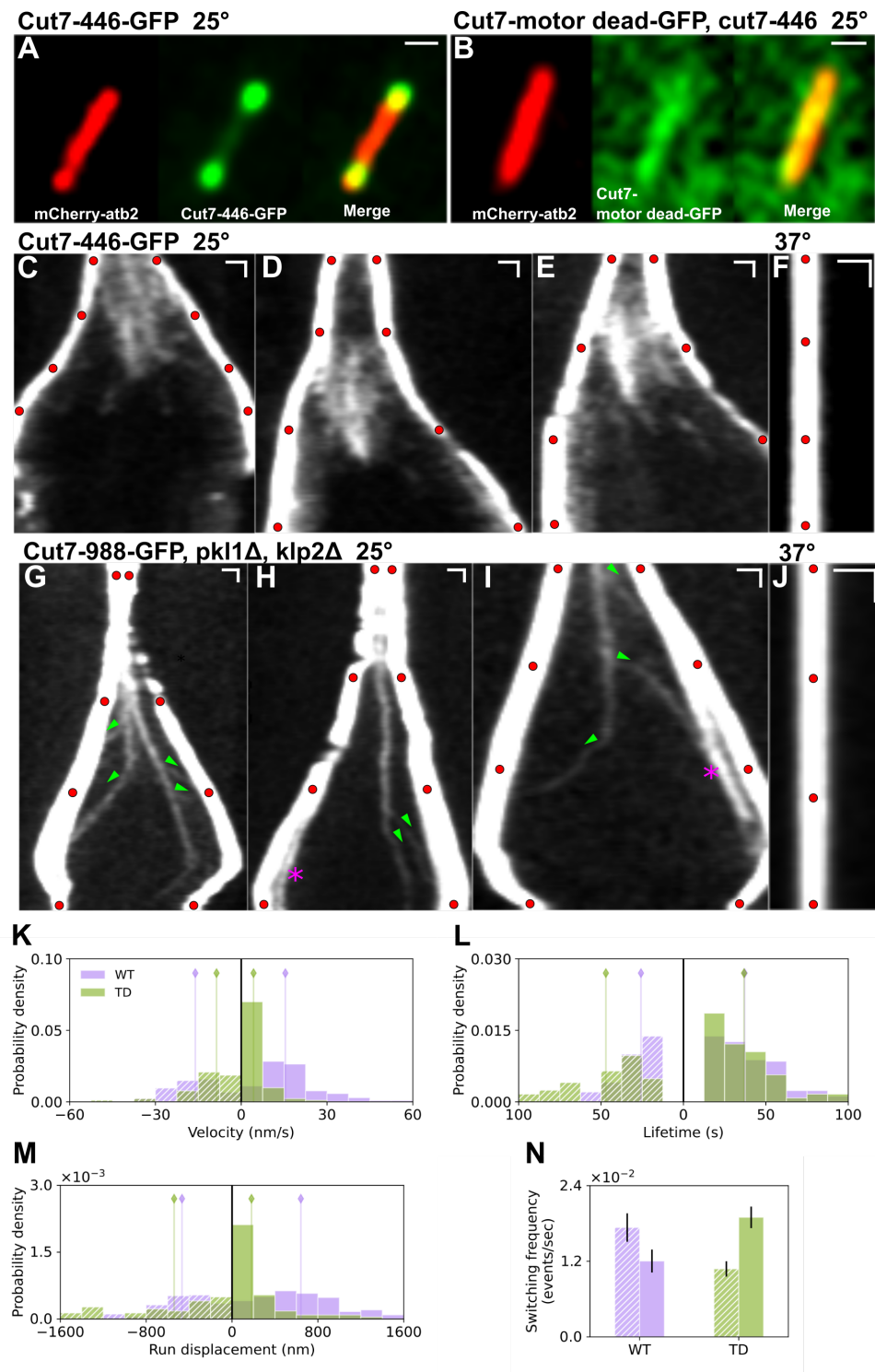


Fig. 5. Mutations in Cut7p alter its motility on bipolar and monopolar spindles. (A-B) Two alleles of Cut7-GFP in *cut7-446* cells at permissive temperature. (A) Dual color image of Cut7-446-GFP (green) on bipolar spindles, demonstrating sparse localization along spindle MTs (red) but normal concentration at the poles (B) Analogous image of Cut7-motor dead-GFP in *cut7-446* cells, showing some spindle binding but no concentration at the poles. (C-F) Kymographs of Cut7-446-GFP. Bipolar spindles (C-E) at permissive temperature show motility similar to *cut7*+ cells while monopolar spindles (F) at restrictive temperature show severely reduced plus-end-directed motility compared to Cut7-GFP. (G-N) Cut7-988-GFP kymographs and data. Removal of the *cut7* tail domain in the absence of *pkl1* and *klp2* leads to reduced motility in bipolar spindles (G-I) and reduced plus-end-directed motility in monopolar spindles (J). SPBs: red circles; Vertical scale bars: 60 s; Horizontal scale bars 1 μ m. (K-N) Comparison of Cut7-GFP and Cut7-988-GFP motility. Values corresponding to poleward-directed movement are on the left side of the axis, anti-poleward-directed movement on the right. Vertical bars represent median of the distribution. For quantification, see table S2. (K) Velocity. (L) Lifetime. (M) Run displacement (N) Switching frequency out of each directed event.

to Cut7-motor-dead-GFP (Fig. 5B, S9). However, Cut7-988-GFP appeared present at lower levels on the interpolar spindle than Cut7-GFP in the same strain background (Fig. 5G-I compared to Fig. 2A-D). One possible explanation of the reduced signal intensity in the spindle midzone could be that Cut7-988-GFP is present at lower levels in the cell or is impaired in microtubule binding. To test this, we compared total intensity of Cut7-GFP and Cut7-988-GFP in line scans along the spindle (Fig. S11). This analysis showed that the integrated intensity is comparable for the two alleles of Cut7 and in fact Cut7-988-GFP showed slightly higher total intensity. Therefore, it appears that the reduced midzone intensity of Cut7-988-GFP likely results from enhanced spindle-pole localization. This suggests that the Cut7 tail may play a role in either plus-end-directed movement or in stabilizing microtubule binding once the motor has reached the midzone. The Cut7-988-GFP that was observed near the midzone in early mitosis lay in faint clusters with reduced movement (Fig. 5G-I, note that the brightness is increased in these figure panels to make the midzone signal visible; for brightness-matched versions of the kymographs see Fig. S12). In late anaphase B, however, similar clusters showed processive movement toward spindle poles (Fig. 5G-I), green arrowheads, S13). The poleward movement typically continued until the signal reached the spindle pole. On monopolar spindles, we observed no plus-end-directed movement (Fig. 5J), consistent with the lethal phenotype of this truncation allele when kinesin-14s are present. Together, these data suggest that Cut7-988-GFP has lost most plus end-directed motility but retains minus end-directed motion.

We quantified Cut7-988-GFP movement and compared it to Cut7-GFP movement on bipolar spindles. Because *cut7-988-GFP* cells appeared to show defects in spindle elongation, we could not easily identify the onset of anaphase B from spindle length changes. Therefore, we examined all directed movement events, independent of stage of mitosis, for both Cut7-GFP and Cut7-988-GFP (Fig. 5K-N, S13, S14A,B, Table S1, S2). The speed of Cut7-988-GFP movement was 2-3 times slower than Cut7-GFP, depending on the direction of movement (Fig. 5K). This is in contrast to human kinesin-5/Eg5, which showed faster motility *in vitro* when the tail was truncated (Bodrug et al., 2020). The lifetime of poleward movement events was nearly two times longer for Cut7-988-GFP than for Cut7-GFP, while the lifetime of anti-poleward movement was comparable (Fig. 5L). As a result, the run displacement of anti-poleward movement was ~ 3 times smaller for Cut7-988-GFP than for Cut7-GFP (Fig. 5M). Consistent with this, the switching rate out of anti-poleward movement was higher than the exit rate from poleward movement for Cut7-988-GFP; this is the reverse of the trend for Cut7-GFP (Fig. 5N). As for the full-length protein, we found no significant correlation between Cut7-GFP spot intensity and particle velocity, lifetime, or run length (Fig. S3, green).

To test whether the minus-end-directed movement we observed in late anaphase was due to Cut7-988-GFP motor activity or to immobile Cut7p on depolymerizing microtubules in the final stages of anaphase, we monitored the time at which the mCherry-tubulin signal in the spindle decreased and the time when Cut7-988-GFP moved poleward. The Cut7p poleward movement typically preceded the microtubule signal decrease (Fig. S14C,D, white arrowheads). Thus, late anaphase Cut7-988-GFP movement is likely minus-end-directed motor activity. At the same stage of anaphase, we also observed bright, fluorescent, relatively immobile clusters of Cut7-988-GFP near the spindle poles (Fig. 5H,I, magenta asterisks).

These results are consistent with the hypothesis that Cut7p bidirectional movement is a result of motor activity, and while MT dynamics may affect its motility it is not the primary cause. The lack of Cut7-988-GFP signal in the spindle midzone could reflect a reduced affinity of the motor for microtubule overlaps, leading to enhanced binding at the pole. Our quantification of Cut7-988-GFP motility suggest that the C-terminal tail beyond amino acid 988 is a determinant of Cut7 motor activity, with a contribution to plus-end-directed movement.

Human Eg5 that replaces Cut7p shows altered localization and motility on fission-yeast spindles

While several mutations to *cut7* appeared to either abolish its movement or favor MT minus-end-directed motility (Fig. 5), we did not identify *cut7* mutants with a greater propensity for plus-end-directed movement. An alternative to study plus-end-directed movement is the human kinesin-5/Eg5, a plus-end-directed motor that has recently been shown to complement *cut7* as the sole kinesin-5 in *S. pombe* (Hwang et al., 2022). Consistent with prior work, we found that cells containing Cut7-GFP or Eg5-GFP expressed at low levels in the *cut7* deletion background were viable (Fig. 6, S15). While Cut7-GFP localized brightly to the spindle poles and more dimly along spindle MTs (Fig. 6A), Eg5-GFP was visible along the spindle with little to no enhancement at the poles (Fig. 6B). To further examine the dynamics of motor localization over time, we compared kymographs of Cut7-GFP and Eg5-GFP on bipolar spindles. The pole localization is noticeable for Cut7-GFP (Fig. 6C), while Eg5-GFP is visible at the spindle poles only rarely and transiently (Fig. 6D). We also observed that in Eg5-GFP cells, initial spindle pole separation often occurred more quickly and was followed in about 20% of cells by spindle shortening events (Fig. 6D). These results suggest that when a purely plus-end-directed kinesin-5 motor is present on the fission yeast spindle, it may be over-active and drive premature spindle elongation followed by shortening.

This relatively uniform Eg5-GFP spindle localization is consistent with our other results suggesting that the bidirectional motility of Cut7 contributes to its pole localization; plus-end-directed Eg5 would therefore not be predicted to accumulate at the spindle poles. In addition, Eg5-GFP fluorescence visible both at the spindle midzone and near spindle poles is consistent with direct motor binding along the length of the spindle. Eg5-GFP may therefore show different localization from Cut7-GFP in part because Cut7p binds directly to γ -tubulin at spindle poles (Olmsted et al., 2014).

DISCUSSION

Cut7-GFP moves bidirectionally on spindle microtubules with regulated motility

Our results show that Cut7p can move in cells toward both plus and minus ends of mitotic microtubules, similar to its bidirectional motility *in vitro* (Edamatsu, 2014; 2016; Britto et al., 2016). After verifying that Cut7-GFP localized on the spindle, as observed in previous work (Fig. 1, Hagan and Yanagida (1992); Drummond and Hagan (1998); Yukawa et al. (2015)), we examined the directional movement of Cut7-GFP on both bipolar and monopolar spindles. Cut7-GFP fluorescent signal moved in both directions along bipolar spindles

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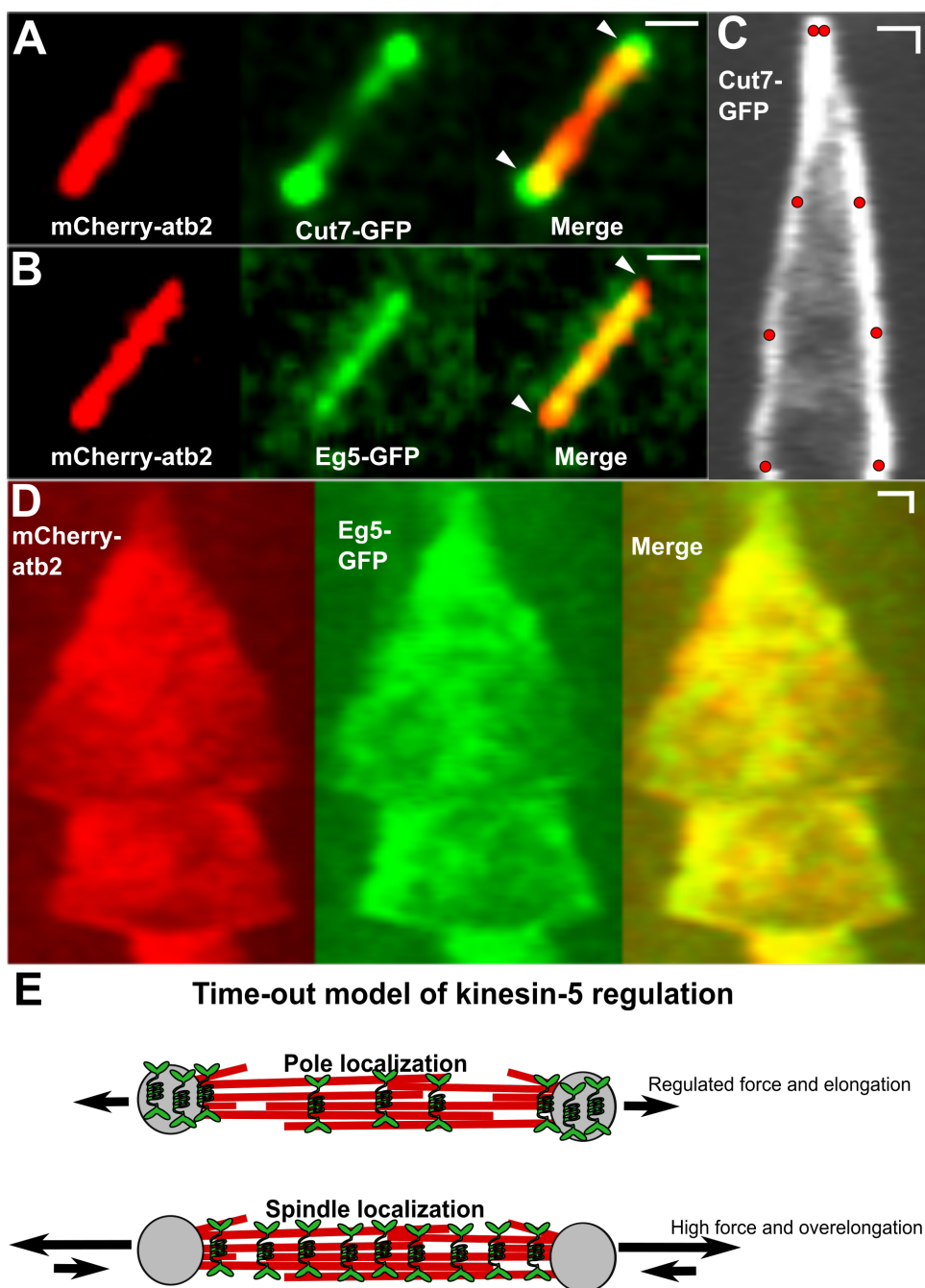


Fig. 6. Eg5-GFP shows altered localization on bipolar spindles. (A-B) Dual color images of bipolar spindles. (A) Cut7-GFP (green) showing typical enrichment at the SPBs (arrowheads) and binding along the MTs (red). (B) SPB enrichment in Eg5-GFP (green) is no longer present (arrowheads) and Eg5-GFP binds uniformly along the spindle. (C-D) Kymographs of bipolar spindles. (C) Cut7-GFP shows consistent SPB enrichment and movement along the interpolar spindle. (D) Eg5-GFP shows reduced SPB enrichment and more uniform localization along spindle microtubules. Spindle shortening events also occur in the Eg5-GFP. SPBs: red circles; Vertical scale bars: 60 s; Horizontal scale bars: 1 μm. (E) Schematic model illustrating the hypothesis that spindle-pole localization may sequester kinesin-5 motors to regulate force generation.

360 (Fig. 2), which is consistent with bidirectional motility but does not definitively demonstrate it because the spindle contains extensive
 361 overlapping antiparallel microtubules. The bidirectional movement also occurred on monopolar spindles that contain microtubules of
 362 only one orientation (Fig. 3, 4). We note that Cut7-GFP movement on bipolar and monopolar spindles was similar, and occurred during
 363 comparable time in mitosis in all cells. The similar motility observed therefore suggests that motor direction is defined by features of the
 364 motors themselves, rather than being strongly influenced by microtubule polarity and parallel versus antiparallel crosslinking. Consistent
 365 with this view, we observed that individual fluorescent spots can change their direction of motion (Fig. 3D-G, S5, S8).

The speed of directed events that we measured was \sim 15–30 nm s $^{-1}$ for both directions of movement (Fig. 2L, 3I, Table S2). In previous experiments with purified Cut7p, surface-bound motors drove microtubule motion in gliding assays at similar speeds, \sim 10–30 nm s $^{-1}$ (Edamatsu, 2014; Britto et al., 2016), although the speed was higher (\sim 200 nm s $^{-1}$) for minus-end-directed movement at low concentration (Britto et al., 2016). Motion of Cut7 on single microtubules occurred at 150–200 nm s $^{-1}$ (Edamatsu, 2016). Thus our velocity measurements are consistent with the slower range of speed measured *in vitro* and the speed of Eg5 *in vitro* (Bodrug et al., 2020).

Our findings suggest that while Cut7p movement has a stochastic component, it is regulated and changes as mitosis progresses. In particular, we found that the activity of Cut7-GFP increased at anaphase B onset on both bipolar and monopolar spindles. We observed a larger percentage of cells that showed Cut7-GFP movement, a greater frequency of directed events, and an increased rate of switching in anaphase B (Fig. 2, 3, 4, table S1). We speculate that phosphoregulation of Cut7p could play a role in this change in activity, as has been previously demonstrated for budding-yeast kinesin-5/Cin8 (Avunie-Masala et al., 2011; Shapira and Gheber, 2016). The *cut7* C-terminal tail contains several conserved phosphorylation sites, including a Cdk consensus site in the BimC box (Heck et al., 1993; Sawin and Mitchison, 1995; Blangy et al., 1995; Drummond and Hagan, 1998; Cahu et al., 2008; Rapley et al., 2008; Akera et al., 2015). In addition, multiple sites in the *cut7* tail have been shown to be phosphorylated in cells by mass spectrometry (Wilson-Grady et al., 2008; Koch et al., 2011; Kettenbach et al., 2015). This makes tail phosphorylation of interest for future study.

These observations of increased Cut7-GFP activity and fluorescence signal in the spindle midzone is consistent with its contribution to anaphase spindle elongation (Yukawa et al. (2019), Fig. 2). The fact that most Cut7-GFP signal disappears from the spindle midzone not long after anaphase B starts is also consistent with previous work showing that kinesin-6/Klp9 is the primary motor driving spindle elongation in *S. pombe* (Fu et al., 2009). By late anaphase, the Cut7-GFP still visible on the spindle appears to be primarily minus end-directed, and thus likely does not contribute significantly to spindle elongation (Fig. 2G). Therefore, it is likely that any contribution of Cut7p to spindle elongation by antiparallel microtubule sliding occurs early in anaphase.

Motor and tail domains both contribute to Cut7 bidirectional motility and pole localization

We find that both the motor and C-terminal tail domains of Cut7 contribute to its bidirectional movement and localization. The tail mutations we examined appeared to limit the motility of Cut7p toward MT plus ends, which suggests that the ability of Cut7p to move toward MT plus ends is more fragile than its minus-end-directed motion. Both a point mutation in the tail region (*cut7-446*) and tail truncation (*cut7-988*) favor the motor's pole localization and appear to reduce its plus-end-directed motility (Fig. 5F,J). The results also point to a role of the C-terminal tail domain as a site for motor regulation. Some properties or modification of the tail may be important for inducing plus end-directed motion of Cut7p.

While wildtype *S. cerevisiae* Cin8 localizes to the minus ends at the SPB of pre-assembled (monopolar) spindles, a mutant lacking the C-terminal tail (but still containing a nuclear localization signal) localizes to the plus ends of monopolar spindles (Shapira et al., 2017). On bipolar spindles, this mutant localizes similarly to a wildtype Cin8, at the spindle poles and the interpolar spindle (Düselder et al., 2015). Therefore, the tail truncation of Cin8 appears to favor its plus-end-directed motility. In contrast, here we show that the Cut7-988-GFP tail deletion mutant compared to full-length shows an increased bias toward the poles of both bipolar and monopolar spindles.

Truncation of the human kinesin-5/Eg5 tail causes it to move more quickly and cluster less *in vitro*, possibly because tail-motor interactions alter the motor conformation to slow ATP hydrolysis (Bodrug et al., 2020). In contrast, we find in cells that the tail-truncated Cut7-988-GFP moves more slowly; however, we also see decreased clustering. Despite these differences, both motors appear to be impaired in sliding MTs since *cut7-988* is lethal when it is the only *cut7* allele present in the *pkl1+ klp2+* background. For Eg5, decreased MT sliding force generation was directly measured *in vitro* (Bodrug et al., 2020), while our results indirectly suggest that Cut7-988p has defects in force generation because it is lethal unless the opposing kinesin-14s are also deleted.

The bidirectional movement of Cut7p that we observed could in principle be driven by tip-associated Cut7p on dynamic spindle MTs. To test this idea, we examined how Cut7-GFP localization and motility changed in cells when MT dynamics or Cut7p motility were perturbed. Deletion of kinesin-8/Klp5 induces elongation of cellular microtubules and decreases their dynamicity (Unsworth et al., 2008). We found that Cut7-GFP motility showed only subtle changes on monopolar spindles in *klp5Δ* cells (Fig. 4), supporting that idea that microtubule dynamics are not the primary driver of the observed Cut7-GFP movement. In contrast, perturbations to Cut7p itself did lead to changes in its localization and motility (Fig. 5). Our observation that Cut7p accumulates poorly at SPBs when its ATPase activity is eliminated (Fig. 5B) provides evidence that Cut7p spindle pole accumulation is at least in part motor-driven. Consistent with this idea, plus-end-directed human Eg5 can replace Cut7p in *S. pombe* (Hwang et al. (2022), Fig. S15), but does not strongly localize to spindle poles (Fig. 6). We conclude that Cut7p motor activity contributes to the directed movement we have described.

Function of fission-yeast versus budding-yeast kinesin-5s

The budding yeast *S. cerevisiae* contains two kinesin-5s, Kip1 and Cin8. Of these, Cin8 appears to be more similar to Cut7p (51.2% BLAST sequence identity versus 35.4% sequence identity with Kip1). Both budding yeast motors are strongly localized to the spindle poles and more weakly at or near the spindle midzone, but neither of them accumulates there until some time into anaphase (Gerson-Gurwitz et al., 2011; Fridman et al., 2013). This pattern of localization is somewhat similar to that of Cut7p in *S. pombe*. However, Cut7p does not concentrate at the midzone in late anaphase, as Cin8p does, probably reflecting the fact that kinesin-6/Klp9 is important for late anaphase B spindle elongation in fission yeast, while the kinesin-5s play this role in budding yeast. Both Cin8 and Cut7p have been shown to be bidirectional *in vitro* (Roostalu et al., 2011; Gerson-Gurwitz et al., 2011; Edamatsu, 2014; 2016; Britto et al., 2016), and Cin8 also moves bidirectionally on spindle MTs *in vivo* (Gerson-Gurwitz et al., 2011; Fridman et al., 2013). Our work thus confirms

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424 that Cut7p moves bidirectionally on the spindle *in vivo* as well. Both motors are important for separation of SPBs to establish a bipolar
425 spindle (Hoyt et al., 1992; Roof et al., 1992; Hagan and Yanagida, 1990; 1992; Tallada et al., 2009), suggesting that both exert plus-end-
426 directed forces early in mitosis. In contrast to Cut7p, Cin8 appears to persist in this role throughout mitosis, while Cut7p passes that role
427 on to a different motor.

428 In *S. cerevisiae*, distinct, processive Cin8 tracks appear common (Gerson-Gurwitz et al., 2011). In *S. pombe*, clear tracks can be
429 identified but appear less common (Fig. 2). Instead, a region of dim Cut7-GFP fluorescence is visible along the spindle. We observed
430 distinct tracks occasionally in cells with Cut7-GFP and frequently in cells with the tail truncated Cut7-988-GFP (Fig. 5G-I). Therefore,
431 the region of dim Cut7-GFP fluorescence on bipolar spindles likely reflects combined movement of many motors.

432 The minus-end-directed motility of Cin8 depends on phosphorylation in the motor domain on Cdk1 sites not conserved in *cut7*
433 (Shapira and Gheber, 2016). By contrast, we find that the Cut7p perturbations we examined decreased or abolished plus-end-directed
434 movement. Our results therefore suggest that Cut7p is more intrinsically minus-end-directed and acquires plus-end directionality by
435 some mechanism that has not yet been clearly identified. This may be due to motor crowding (Britto et al., 2016), phosphorylation
436 analogous to the regulation of Cin8, or some other mechanism.

437 Function of Cut7p bidirectional motility and pole localization

438 It is well established that kinesin-5s are able to crosslink and slide antiparallel MTs to separate spindle poles, a function which depends
439 on plus-end-directed motility of kinesin-5 tetramers. Our work further supports the importance of kinesin-5 plus-end-directed motility,
440 because the temperature-sensitive *cut7-446* and *cut7-988* alleles that show either no motility or reduced plus-end-directed events on
441 monopolar spindles at 37°C are unable to create a bipolar spindle (Fig. 5F, J). Consistent with this, *cut7-motor-dead*, which does not
442 move on bipolar spindles (Fig. S9), is lethal when it is the sole *cut7* allele in the cell (Akera et al., 2015). Given the importance of
443 plus-end-directed motility, the importance of bidirectional motility/trafficking and pole localization has been less clear.

444 In previous work, we used a computational model of *S. pombe* spindle assembly to propose that localization of kinesin-5/Cut7 near
445 the spindle poles may be required for proper spindle-pole separation (Blackwell et al., 2017; Edelmaier et al., 2020). This was based
446 on model simulations in which a processive, purely plus-end-directed kinesin-5 localized far from the SPBs in monopolar spindles and
447 failed to separate spindle poles. Experimental work on budding-yeast Cin8 reached a similar conclusion (Shapira et al., 2017). However,
448 the experimental results we present here appear to rule out that hypothesis. Importantly, human Eg5 can replace *cut7* and assemble a
449 bipolar spindle (Hwang et al., 2022) despite being a plus-end-directed kinesin-5 and not significantly localizing to *S. pombe* spindle poles
450 (Fig. 6). Therefore, minus-end-directed motility is not required for bipolar spindle assembly in fission yeast. How to reconcile this with
451 the computational modeling results is an interesting question for future work. We speculate that perhaps Eg5 has higher turnover that
452 allows it to unbind and avoid becoming localized primarily at the plus-ends of spindle microtubules in early mitosis.

453 We observed that replacement of Cut7p with low-level expression of Eg5 in *S. pombe* can lead to faster initial separation of spindle
454 poles, a longer spindle, and transient spindle shortening (Fig. 6D). Additionally, recent work found that high-level expression of either
455 human Eg5 or Cut7p in *S. pombe* was lethal with over-elongated spindles (Hwang et al., 2022). Together, these results suggest that a
456 highly expressed or purely plus-end-directed kinesin-5 that localizes along spindle MTs may produce excessive sliding force that over-
457 elongates the spindle. Therefore, we propose a new hypothesis that spindle-pole localization of kinesin-5 may sequester the motor in a
458 “time-out” at a location where its sliding activity is low (Fig. 6E). In this view, the spindle-pole localization may decrease motor activity.
459 This model will be of interest to test in future work.

460 MATERIALS AND METHODS

461 Strain construction by crossing

462 Cells were cultured using standard techniques (Moreno et al., 1991). Existing strains (table S4) were crossed and the desired phenotype
463 isolated using random spore analysis. The *cut11-7* and *cut7-446* mutations were identified by replica plating colonies from the spore
464 analysis onto YE5S plus phloxin B agar plates. After 2–3 d at 37°C, dark-pink colonies were identified as positive for the mutations. All
465 other genes were identified using EMM plates without the relevant supplement for auxotrophic mutants or YE5S plates with the relevant
466 antibiotic. The fluorescent label for microtubules was obtained by expressing an mCherry- α -tubulin-chimera at a low level (~10% wild
467 type α -tubulin), as used previously (Yamagishi et al., 2012; Gergely et al., 2016; Blackwell et al., 2017). This low-level labeling reduces
468 possible tag-related perturbations to microtubule dynamics. Cells with kinesin-5 mutation or deletion are susceptible to chromosome
469 missegregation and other mitotic abnormalities. To avoid problems resulting from these abnormalities, cells for each experiment were
470 stored at 4°C and reisolated from frozen stocks every 2–4 weeks.

471 Plasmid and strain construction by molecular biology

472 Oligonucleotide primers were purchased from Integrated DNA Technologies (Coralville, IA). Restriction enzymes and Phusion HF
473 DNA polymerase were purchased from New England Biolabs (Ipswich, MA). DNA was prepared using Qiaprep Spin Miniprep Kit
474 and polymerase chain reaction (PCR) products were purified using Qiaquick PCR Purification Kit, both from Qiagen (Germantown,
475 MD). *S. pombe* genomic DNA was prepared using YeaStar Genomic DNA Kit from Zymo Research (Irvine, CA). DNA sequencing was
476 performed by Quintarabio (Hayward, CA). DNA concentration was determined using a Thermo Scientific Nanodrop 2000. Strains were
477 verified by PCR and sequence analysis of genomic DNA. At least two different transformant strains were analyzed in experiments.

Strain construction for Cut7-photoactivatable(PA)-GFP

PA-sfGFP (Addgene 54579) was amplified by PCR using oligonucleotide primers Pac1-paGFP-F (GGGTTAACGTGAG-CAAGGGCGAGGAG) and Asc1-paGFP-R (AGTGGCGCGCCCTACTTGTACAGCTCGTCCATGCC) and the product was cloned into pFA6a-GFP(S65T)-kanmx6 (Addgene 39292) digested with Pac1/Asc1. The resulting plasmid was amplified by PCR using oligonucleotide primers Cut7-Cterm-pFA6a-GFP/paGFP-F (AATTCAAGAACTAGTCTTGGAGTAGCAGAAGT-GCCTATCCAAAATGAAACGACGGATCCCCGGTTAATTAA) and Cut7-Cterm-pFA6a-GFP/paGFP-R (GATGTAATACATTCTATTGTATTCGTCCATTAAGTATAAAATCGTCAGAATTGAGCTCGTTAAC) and the product was used to transform *S. pombe* using a lithium acetate method (Okazaki et al., 1990).

Strain construction for Cut7-446-GFP and Cut7-988-GFP

For *cut7-446-GFP*, strain MB921 was transformed with PCR product obtained using plasmid pFA6a-GFP(S65T)-kanmx6 (Addgene 39292) and oligonucleotide primers Cut7-Cterm-pFA6a-GFP/paGFP-F and Cut7-Cterm-pFA6a-GFP/paGFP-R. For *Cut7-988-GFP*, strain MB1147 was transformed with PCR product obtained using plasmid pFA6a-GFP(S65T)-kanmx6 (Addgene 39292) and oligonucleotide primers Cut7-988-Cterm-pFA6a-GFP-F (TTAAAGGAAACGACATCACTGCTAATCATACTAATGAATTACTGGTTAG-GAGATGAACGGATCCCCGGTTAATTAA) and Cut7-Cterm-pFA6a-GFP/paGFP-R. The choice to truncate at amino acid 988 was based on coiled-coil prediction (Lupas et al. (1991), Fig. S10). All of these *cut7* alleles contain the identical single GFP molecule separated from the *cut7* open reading frame by a six amino acid linker (RIPGLI).

Strains expressed from heterologous promoter

The genes for *cut7*, *cut7-motor-dead*, and Eg5 were cloned using PCR into plasmid 462 (Addgene 89065) containing a Z3EV promoter (Ohira et al., 2017) modified to contain Asc1/Srf1 sites just upstream of the GFP gene. The motor dead mutation (G164A-K165A-T166A) was cloned by PCR from strain PB951 (MB947, Akera et al. (2015)) into the 462-cut7 plasmid. These alleles all contain the identical single GFP molecule (as for the *cut7* alleles discussed above), but for these *cut7* alleles the linker is three amino acids (SRA). For the Eg5 allele expressed under the heterologous promoter, there is no linker before the GFP. These plasmids were cut at AatII and integrated at the *his7* gene in Strain FY31411 (yFS949, MB1138). The strain was crossed to produce strain MB1164. Estradiol was not used in our experiments; rather *cut7* or Eg5 were expressed at a constitutive lower level.

Live-cell imaging

All microscopy images and related datasets were obtained from confocal microscopy using live-cell preparation. Cells were grown at 25°C on YE5S plates for 48 hours and restreaked every 12 hours. A small volume of exponentially growing cells was removed from the petri dish and placed in 10 µL of EMM. EMM was filtered with a 0.2 µM cellulose acetate filter to reduce background fluorescence. EMM was placed onto 22 × 60-mm coverslips coated with 8 µL of lectins from *Bandeiraea simplicifolia* (Sigma-Aldrich). These coverslips were pre-equilibrated to the appropriate temperature of either 25 or 37°C. Bipolar spindles were imaged at 25°C and monopolar spindles were imaged at 37°C. To obtain sufficient monopolar spindles, *cut11-7* cells were placed at 37°C for 2-4 hours and then imaged at restrictive temperature. Cells were transferred from a 37°C incubator to the pre-warmed microscope in less than 30 seconds to prevent monopolar spindles cooling down and possibly becoming bipolar. Temperature was maintained with ±0.1°C precision using a CherryTemp temperature controller (Cherry Biotech, Rennes, France). Spinning-disk confocal microscopy was performed on a Nikon Eclipse Ti microscope described previously (Gergely et al., 2016; Blackwell et al., 2017; Edelmaier et al., 2020). Time-lapse image stacks were obtained using the EM gain laser settings on the Nikon illumination system and number of Z-planes described previously (Edelmaier et al., 2020).

PA-GFP and fluorescence photobleaching experiments

PA-GFP experiments used a 405 nm, 50 mW coherent Obis laser (Santa Clara, CA). Before photoactivation, 4 image stacks with illumination from the 561 nm and 488 nm lasers were obtained to record the location of the microtubules and confirm that no GFP signal existed prior to activation. Photoactivation was then performed on either a monopolar microtubule bundle or near the spindle pole with the laser at 10% power and an exposure time between 15-1000 ms. After photoactivation, time-lapse image stacks were obtained continuously using the 488 nm illumination laser. For fluorescence photobleaching experiments, the same 405 nm laser was used. Before photobleaching, 4 image stacks with illumination from the 488 nm laser were obtained to record the location of the spindle poles and Cut7-GFP. Photobleaching was then performed on the area of interest with the laser at 50-100% power and an exposure time of 100 ms. After photobleaching, signal was measured with time-lapse image stacks obtained continuously using the 488 nm illumination laser.

Determination of spindle length for pre-anaphase and anaphase B

Ten monopolar spindles each for *klp5+* and *klp5Δ* cells were identified which underwent rapid elongation after a period of a constant, stable length. The average length at this transition for *klp5+* spindles was 2.40 µm, and for *klp5Δ* spindles was 5.16 µm.

Dataset organization and kymograph creation

For all bipolar tracked data in this study, strains with Cut7-GFP or Cut7-3GFP, and *cut11+* or *cut11-7* were pooled to create the kymographs and associated datasets displayed in figures 2-5. In addition, PA-GFP and fluorescence photobleaching experiments were pooled

530 with all non-PA-GFP and unbleached experiments to create datasets for figures 2-3. The kymographs used for the quantification in
531 figure 2 were compiled from strains MB951, MB1030, MB1032, MB1062, MB1064, MB1077, and MB1085 (table S4). The kymo-
532 graphs quantified for figure 3 were compiled from strains MB951 and MB1088 (table S4). The monopolar spindles analyzed in figure 4
533 were compiled from strain MB1134. The bipolar spindles analyzed in figure 5 were compiled from strains MB1131 and MB1155 and
534 compared with the pooled data from figure 2.

535 All confocal images were transferred as Nikon nd2 files into the Fiji version of ImageJ (National Institutes of Health, Bethesda, MD)
536 and displayed as pixel-interpolated, maximum-intensity projections. The FIJI plugin StackReg was utilized to align the data using the
537 Transformation-Rigid Body option. The Image J segmented line tool was used to draw a straight line through the aligned spindle data.
538 Finally, the Image J Reslice function with an output spacing of 1 μm was used to create the kymographs shown.

539 **Kymograph preparation and analysis for Cut7-motor-dead**

540 To prepare the Cut7-motor-dead-GFP kymographs (Fig. S9), we identified the spindle axis direction and the centrosome location using the
541 Toolkit for Automated Microtubule Tracking (TAMiT, manuscript in preparation, <https://github.com/Betterton-Lab/TAMiT.git>). Based
542 on spindle length and position fit by TAMiT, we defined and aligned a constant-length axis along the spindle using the red channel in every
543 frame. We then used Matlab to interpolate the pixel intensity along the defined axis to make kymographs for both channels (microtubule
544 and Cut7-motor-dead-GFP, Fig. S9A,B). The brightness and intensity of each channel of the kymograph are adjusted independently in
545 FIJI before merging.

546 To measure the GFP intensity distribution along the spindle axis for both Cut7-GFP and Cut7-motor-dead-GFP (Fig. S9C-F), we
547 identified slices from kymographs corresponding to spindle length of 1, 2, 3, and 4 μm , $\pm 0.1 \mu\text{m}$ around this length. The Cut7-GFP
548 kymograph line-scans were retrieved from 29 Cut7-GFP kymographs, and from 7 Cut7-motor-dead-GFP kymographs. All individual
549 line scans were aligned based on one of the spindle-pole positions (chosen to be the pole with brighter intensity for Cut7-GFP imaging
550 and chosen at random for Cut7-motor-dead-GFP imaging). We then averaged all line scans for the relevant spindle length and plotted the
551 average *cut7* intensity distribution along the spindle axis.

552 **Total intensity analysis for Cut7-GFP and Cut7-988-GFP**

553 To quantify the total GFP intensity along the spindle axis for both Cut7-GFP and Cut7-988-GFP (Fig. S11), we identified slices from
554 kymographs corresponding to spindle length of 2.5-5.5 μm . The Cut7-GFP kymograph line-scans were retrieved from 1162 frames
555 collected from 29 Cut7-GFP cells, and from 1169 frames collected from 40 Cut7-988-GFP cells. All individual line scans were aligned
556 based on one of the spindle-pole positions (chosen to be the pole with brighter intensity). We then summed all line scans to determine
557 the total Cut7-GFP intensity along the spindle axis.

558 **Track and SPB annotation**

559 Annotations were made directly on the kymograph. We created in-house MATLAB software that allowed a user to interactively draw
560 Cut7p tracks (Fig. S2, S5, S8, S13). Some of these tracks were identified as corresponding to spindle-pole-localized Cut7-GFP, while
561 other tracks were used for motility analysis. For bipolar spindle kymographs, both spindle poles were identified by areas of highest
562 brightness throughout the kymograph. Two SPB lines were traced over those areas. Clusters of Cut7-GFP moving along spindle micro-
563 tubules were then traced. The nearest SPB was used as a reference for describing poleward or anti-poleward movement; events that
564 showed movement all the way from one pole to the other were excluded from analysis due to ambiguity in identifying the nearest SPB.
565 For monopolar spindle kymographs, the SPB was identified by the area of highest brightness throughout the kymograph, and an SPB
566 line was traced. Clusters of Cut7-GFP moving poleward or antipoleward were then traced.

567 **Track merging and event identification**

568 We started by merging tracks that were deemed to belong to the same global track. For example, if track 2 began within some time
569 interval of track 1 ending, and the start position of track 2 was close to the end position of track 1, then we merged the tracks (Fig. S1).
570 After a phase of merging, we identified a number of tracks with both poleward and anti-poleward motion. The tracks were then analyzed
571 by splitting them into poleward, anti-poleward, and paused segments or events. Event identification was based on the average local
572 velocity along the track. A cutoff value of 3 nm s^{-1} was used to distinguish directed movement from pausing. Points with velocity less
573 than the negative of the cutoff are assigned as poleward, while points with velocity greater than the cutoff are assigned as antipoleward.
574 The remaining points were assigned as paused.

575 **Track analysis quantification**

576 Tracks were analyzed with reference to the position of the SPB. To do that, we assigned each track to its nearest SPB. For monopolar
577 spindles, there was only one pole and assignment was straightforward. For bipolar spindles with two SPBs, we picked the SPB that
578 was closest to the track start position. Tracks that moved all the way across the bipolar spindle from one pole to another were excluded
579 from this analysis. Distance and velocity were measured relative to the closest SPB. For each kymograph, the frequency of directional
580 events was calculated by counting the number of poleward and anti-poleward tracks and dividing by the total time of the kymograph. The
581 switching frequency was calculated by counting the number of switches out of a state (poleward and anti-poleward movement) divided
582 by the total time of all tracks in that state. For example, the switching frequency for exiting the poleward-moving state was calculated by
583 dividing the number of transitions out of the poleward state by the total time of all poleward tracks.

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The authors report no competing interests.	591
Contribution	592
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Data availability	598
All data and computer code used in this study are available upon request.	599
Supplementary	600
See supplementary figures and tables in the attached file.	601
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