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Accessibility
Microtubule sliding activity of a kinesin-8 promotes spindle assembly and spindle length control

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Abstract

Molecular motors play critical roles in the formation of mitotic spindles, either through controlling the stability of individual microtubules, or by cross-linking and sliding microtubule arrays. Kinesin-8 motors are best known for their regulatory roles in controlling microtubule dynamics. They contain microtubule-destabilizing activities, and restrict spindle length in a wide variety of cell types and organisms. Here, we report for the first time on an anti-parallel microtubule-sliding activity of the budding yeast kinesin-8, Kip3. The in vivo importance of this sliding activity was established through the identification of complementary Kip3 mutants that separate the sliding activity and microtubule destabilizing activity. In conjunction with kinesin-5/Cin8, the sliding activity of Kip3 promotes bipolar spindle assembly and the maintenance of genome stability. We propose a “slide-disassemble” model where Kip3’s sliding and destabilizing activity balance during pre-anaphase. This facilitates normal spindle assembly. However, Kip3’s destabilizing activity dominates in late anaphase, inhibiting spindle elongation and ultimately promoting spindle disassembly.

INTRODUCTION

During cell division, the spindle apparatus drives the spatial separation of sister chromatids over a distance up to tens of microns. Spindle length therefore influences the fidelity of chromosome segregation1, 2. The functional and dynamic properties of mitotic spindles are also reflected by characteristic spindle lengths3. Gene deletion and RNAi-based approaches in the past have successfully identified numerous factors that participate in spindle length regulation. These include microtubule regulators that affect the nucleation and assembly of individual microtubules, force-generating motors that sort and align microtubules, as well as factors that maintain chromosome cohesion and the structure of kinetochores3-6. Cooperation and antagonism between different regulators have been revealed by...
combinatorial analysis of gene deletions or gene knockdowns. However, even a single gene product can have complex, potentially counteracting roles in spindle length control, which could be obscured in the analysis of simple gene deletions or knockdowns. For this reason, the functional heterogeneity of individual regulators and its relevance to spindle length control are not well understood.

Here we combine in vitro assays with genetic analysis to define counterbalancing activities of a conserved regulator of spindle length, the budding yeast kinesin-8 motor, Kip3. Our analysis illustrates the importance of considering this level of complexity for understanding sophisticated cellular machineries such as the mitotic spindle. Kinesin-8 motors, best known as microtubule destabilizing factors, restrict metaphase spindle length in a variety of organisms and cell types. However, Kip3 seems to be an exception because kip3Δ cells do not display altered pre-anaphase (metaphase-like) spindle length. This is surprising because Kip3 clearly regulates the length of individual microtubules during pre-anaphase. By contrast, Kip3 restricts spindle length at the end of anaphase when spindles fully extend. Thus, the impact of Kip3 on the spindle varies during mitosis in a way that is not simply predicted by its known biochemical properties. Because of our recent work characterizing a non-motor microtubule binding activity in the Kip3 tail domain, we considered the possibility that Kip3 slides anti-parallel microtubules, an activity that could potentially increase spindle length and counterbalance its activity in shortening spindles by promoting microtubule disassembly.

RESULTS

Kip3 slides apart anti-parallel microtubules

A fluorescence microscopy-based assay was used to determine if Kip3 cross-links and slides microtubules. Recombinant Kip3 and a tail-truncation (Kip3 ΔT-LZ) were purified from yeast (Fig. 1a and Supplementary Fig. S1). Kip3 or Kip3ΔT-LZ was pre-bound to track microtubules (green) that were immobilized on a coverslip. Cargo microtubules (red) were then flowed into the assay chamber. We used Taxol-GMPCPP double stabilized cargo microtubules in these experiments to prevent the Kip3-mediated microtubule depolymerization. We found that full-length Kip3 induced the alignment of cargo microtubules with track microtubules but Kip3ΔT-LZ did not (Fig. 1b). Meanwhile, Kip3 and Kip3ΔT-LZ bound the track microtubules equivalently in this assay (Supplementary Fig. S2). Thus, Kip3 contains a microtubule cross-linking activity that depends on its tail domain. This tail domain contains a secondary microtubule-binding site, in addition to the primary binding site on the motor domain.

Time-lapse imaging revealed that Kip3 drives two types of movements of cargo microtubules along track microtubules: sliding and “tug-of-war” (Supplementary Video 1 and 2). In either case, we often observed an accumulation of Kip3 on one end of cargo microtubules (Fig. 1c). This accumulation likely, though not exclusively, resulted from motors that walked along cargo microtubules and accumulated at the plus ends. Although Kip3 was pre-bound to track microtubules prior to initiating this sliding assay, Kip3 could detach from the track microtubule during the assay, rebind to cargo microtubules, and then walk to their plus ends.

Experiments with polarity-marked microtubules revealed a correlation between sliding and microtubule orientation. When the cargo and track microtubules were in a parallel orientation, Kip3 primarily induced “tug-of-war” movements (Fig. 1d). This is expected if multiple motors bind parallel microtubules with opposite orientations: some Kip3 molecules with the motor domain on the track microtubule, transporting the cargo in one direction, and some Kip3 molecules with the motor domain on the cargo microtubules mediating transport.
in the opposite direction. In this circumstance, the forces will not always be balanced, and some persistent sliding movements were, as expected, observed, but at a low frequency (Supplementary Fig. S3 and S4). By contrast, Kip3 always induced sliding when the microtubules were in an anti-parallel orientation, with the microtubule minus end as the leading end (Fig. 1e, Supplementary Fig. S3, and S4). This is consistent with the well-characterized plus end-directed motility of Kip3\(^{10, 11}\). Moreover, although the majority of cargo microtubules were aligned with track microtubules, occasionally we also found unaligned cargo microtubules (white arrow heads in Fig. 1f) resulting from non-specific attachment of Kip3 to the coverslip, as occurs in standard microtubule gliding assays. We measured the gliding velocity of the unaligned microtubules and it is similar to the sliding velocity of aligned cargo microtubules (Fig. 1f). This suggests a model that when Kip3 is sliding microtubules, the motor domain of dimeric Kip3 moves along one microtubule and the tail domain remains in mostly static contact with the other microtubule. This is similar to dimeric kinesin-14s\(^{24, 25}\) but contrasts with tetrameric kinesin-5s. Kinesin-5s slide anti-parallel microtubules at twice the velocity that they mediate gliding of unaligned microtubules. This is because of the bipolar tetrameric structure of kinesin-5s that enables their motor domains to walk simultaneously on the track and cargo microtubules\(^{26, 27}\). In summary, we have identified an anti-parallel microtubule sliding activity of kinesin-8/Kip3 that could potentially promote spindle assembly and increase the length of mitotic spindles.

**Kip3-mediated sliding of dynamic microtubules**

In addition to the above-described sliding of chemically stabilized microtubules, we found that Kip3 promotes anti-parallel sliding of dynamic microtubules. This was established using recently developed methods using micropatterning technology to control the geometry of microtubule seeds, from which dynamic microtubules were polymerized\(^{28}\). In this assay, dynamic microtubules were polymerized with plus ends growing away from micropatterned sites positioned with 20 \(\mu\)m spacing (Fig. 2a). In the presence of Kip3, microtubules growing from adjacent micropatterned sites formed bundles that subsequently underwent buckling, which was not observed under control conditions (Fig. 2b, Supplementary Video 3 and 4). Because the microtubule seeds were adsorbed on the micropatterned sites, Kip3-dependent sliding was manifest as buckling of microtubule bundles. The degree of buckling was quantified by measuring the deflection of the bundle away from the axis connecting the two asters 30 minutes after the initiation of microtubule polymerization (Figure 2c and Supplementary Fig. S5).

To more faithfully mimic the organization of spindle microtubules, we determined whether Kip3 is able to induce sliding of anti-parallel microtubules bundled by Ase1, the conserved microtubule binding protein that organizes the spindle midzone\(^{29, 30}\). As expected, Ase1 increased the frequency of anti-parallel microtubule bundles from 16% to 60% and most of the bundles formed in the presence of Ase1 were straight (Fig. 2b, 2c, and Supplementary Move S5). Strikingly, when both Kip3 and Ase1 were added together, stable microtubule bundles were formed that steadily elongated and underwent marked buckling (Fig. 2b, 2c, and Supplementary Video 6). Whereas curved bundles were transiently slid in the presence of Kip3 (Supplementary Video 4), the collaboration between Ase1 and Kip3 led to the formation of stable bundles in which microtubule sliding persisted, leading to greater curvatures (Supplementary Video 6). Together, these data demonstrate that Kip3 mediates anti-parallel sliding of its natural substrate: dynamic microtubules cross-linked by Ase1. This supports the conclusion that Kip3 has the ability to promote spindle assembly and increase spindle length.
Identification of a depolymerizing-proficient but sliding-deficient variant of Kip3

Previous genetic and biochemical evidence supports the idea that Kip3 promotes the shortening of microtubules through its microtubule depolymerizing activity\textsuperscript{10, 11, 14}. The microtubule depolymerizing activity is assayed on GMPCPP-stabilized microtubules and has been proposed to reflect the ability to remove the GTP-cap on dynamic microtubules and trigger catastrophe. Whether kinesin-8s other than Kip3 are microtubule depolymerases, however, remains a subject of debate\textsuperscript{15}. Other mechanisms, for example, slowing microtubule growth, have been proposed to explain how these kinesin-8 motors shorten microtubules\textsuperscript{15}. To simplify, we will refer below either to Kip3’s “destabilizing activity” or “depolymerase activity”, acknowledging that the mechanisms for destabilizing dynamic microtubules by kinesin-8s remains under investigation.

In principle, Kip3’s depolymerizing activity and sliding activity could have opposite effects on spindle length control. Such opposing effects could have obscured a role for Kip3 in spindle assembly during previous analysis of kip3\textsuperscript{-Δ} strains. To address this issue, we searched for mutants that would separate these biochemical activities. A candidate mutant that would be deficient for sliding activity but proficient for depolymerization was already in hand. We previously showed that the Kip3 tail domain concentrates Kip3 on the microtubule plus end, and increases the depolymerase activity of Kip3 by approximately two-fold\textsuperscript{21}. This effect is significant because a tail truncation mutant produces a near-null phenotype in cells. However, normal microtubule destabilizing function can be restored by doubling the gene dosage of the tail truncation mutant \textit{in vivo} or by doubling the concentration of the tail truncation mutants \textit{in vitro}\textsuperscript{21}. As shown in Fig. 1b, this tail truncation construct is deficient for microtubule cross-linking and sliding. Thus, this tail-truncation Kip3 mutant is depolymerase proficient but sliding deficient.

Characterizing a sliding-proficient but depolymerizing-deficient variant of Kip3

A mutant that was proficient for microtubule sliding but deficient for microtubule depolymerase activity came out of our analysis of kinesin-8 family specific amino acid sequences. All kinesin motor families share high sequence homology in the motor domain (also called “head”). However, many kinesin families have identifiable family-specific features, one of which is a family-specific neck. Kinesin-8s are one of the kinesin families that appear to have a family-specific neck sequence\textsuperscript{31}. The neck in kinesin-8s is located at the C-terminus of the motor domain and is predicted to form a coiled-coil that mediates dimerization of the motor. To determine if the kinesin-8-specific neck has a role in depolymerase activity, we replaced the Kip3 neck with a leucine zipper motif that maintains the dimeric state of the motor (Supplementary Fig. S1). First, we verified that this Kip3 variant, hereafter referred to as Kip3-CC (Fig. 1a) maintains comparable sliding activity to that of Kip3 (Fig. 3a-c). However, Kip3-CC was markedly deficient in depolymerase activity (~10% of the full-length protein, Fig. 3d). Because Kip3 depolymerizes GMPCPP-stabilized microtubules in a length-dependent manner\textsuperscript{11}, we selected GMPCPP microtubules with a defined range of lengths (3-5 μm) for quantifying depolymerization rates. Moreover, the effects of Kip3 and Kip3-CC on dynamic microtubules assembled \textit{in vitro} were also analyzed. Consistent with another report\textsuperscript{14}, we found that Kip3 promoted microtubule catastrophe. By contrast, Kip3-CC failed to promote microtubule catastrophe, under conditions where the flux of Kip3 and Kip3-CC were comparable (Fig. 3e and Supplementary Fig. S6). Furthermore, Kip3-CC failed to promote the catastrophe of astral microtubules in yeasts, whereas Kip3 increased the catastrophe frequency (Supplementary Table 1). Together, these data suggest Kip3-CC is deficient in promoting microtubule disassembly.
The reduced microtubule destabilizing activity of Kip3-CC could be due to a defect in motility that causes a reduced flux of motors to the microtubule plus end. Alternatively, the reduced depolymerase activity could reflect an intrinsically reduced capacity of Kip3-CC in removing tubulin dimers from the plus end. To distinguish between these two mechanisms, Halo-tagged Kip3 and Kip3-CC were labeled with tetramethylrhodamine (TMR) and their motilities were characterized by single molecule imaging (Fig. 4a). The velocity of Kip3-CC was not significantly different from that of Kip3. We did observe that the average run length of Kip3-CC is reduced by 54% relative to Kip3 (Fig. 4b), however, Kip3-CC is still a highly processive motor with an average run length exceeding 5 microns, longer than almost all microtubules in budding yeast. For microtubules of comparable length to those used to assay depolymerase activity (3–5 μm), equal concentrations of Kip3 and Kip3-CC gave, as expected, similar flux to the plus end (32 Vs 31 min⁻¹), but Kip3-CC was nevertheless markedly deficient for depolymerase activity (Fig. 3d). Thus, the substitution of the Kip3 neck with a leucine zipper motif results in a mutant that markedly reduces microtubule destabilizing activity, but largely maintains motility and sliding activity.

Spindle length is regulated by a balance between the microtubule sliding and destabilizing activity of Kip3

As a first step towards determining whether the sliding activity of Kip3 has a role in spindle length control, we characterized the effects of overexpressing Kip3-CC in cells lacking the endogenous KIP3 gene. Overexpression of Kip3-CC using a pGAL1 expression system caused an apparent increase in spindle length in cycling cells (Fig. 5a). To exclude the possibility that this effect was merely due to altered cell cycles, cells were synchronized in S phase by hydroxyurea. In this case, overexpression of Kip3-CC caused a ~50% increase in spindle length relative to the control (Fig. 5b). This effect is highly significant and comparable to the overexpression of Cin8, the major kinesin-5 sliding motor in budding yeast. Several experiments strengthen the conclusion that it is the sliding activity of Kip3-CC that causes the increase in spindle length (Fig. 5b). First, overexpressing Kp3ΔT-LZ, which lacks microtubule sliding activity, did not increase spindle length. Second, although Kip3-CC overexpression increased spindle length, it did not affect astral microtubule length, suggesting the increase in spindle length does not result simply from an increase in the length of individual microtubules. Third, a Kip3 rigor mutant which cross-links microtubules but does not walk on or slide apart microtubules, did not cause an increase in spindle length, excluding an explanation for the increase in spindle length involving passive cross-linking of microtubules by Kip3 that would increase the sliding effectiveness of other outward sliding motors. This type of mechanism was previously suggested to explain the role of kinesin-14/Kar3 in spindle length control. Fourth, the localization of Ase1 to spindles was significantly reduced when Kip3-CC was overexpressed (Supplementary Fig. S7a and S7b). Ase1/PRC1 is a spindle midzone protein that preferentially binds to anti-parallel microtubules. The reduced localization of Ase1 to spindles suggests a decrease in spindle microtubule overlap, supporting the conclusion that the increase in spindle length results from sliding driven by Kip3-CC. Finally, although we cannot completely exclude the possibility that Kip3-CC indirectly increases spindle length by cargo transport, the midspindle localization of Stu2, the only known Kip3 cargo, was not measurably affected by overexpressing Kip3-CC (Supplementary Fig. S7c and S7d). Together, these data suggest that Kip3-CC increases spindle length because of its microtubule sliding activity.

Next, we attempted to dissect the individual contributions of the sliding and depolymerizing activities of Kip3 to spindle length control. The KIP3 locus was replaced with either kip3-CC (encoding a sliding proficient but depolymerizing deficient motor) or two copies of kip3ΔT-LZ (encoding a depolymerizing proficient but sliding deficient motor). As discussed above, 2x Kip3ΔT-LZ is comparable to Kip3 for its effects on microtubule depolymerization.
in vitro, and confers similar benomyl resistance in vivo\textsuperscript{21}. We found pre-anaphase spindle length was indistinguishable for wild-type and \textit{kip3} \textsuperscript{Δ} cells, consistent with previous reports\textsuperscript{20}. By contrast, pre-anaphase spindles were longer in \textit{kip3-CC} cells, but shorter in 2x \textit{kip3} \textit{Δ-T-LZ} cells relative to the wild-type controls (Fig. 5c). By the end of anaphase, the spindles in \textit{kip3-CC} cells hyper-elongated to an even greater degree than occurs in \textit{kip3} \textit{Δ} cells. 2x \textit{kip3} \textit{Δ-T-LZ} cells exhibited a reciprocal effect where the spindles break down prematurely prior to separation of chromosomes at a spatial maximum (Fig. 5d). We verified that the steady-state protein level of \textit{Kip3-CC} was indistinguishable from that of the wild-type protein, and that doubling of the gene dosage of \textit{kip3} \textit{Δ-T-LZ} resulted in the approximate doubling of \textit{Kip3} \textit{Δ-T-LZ} protein expression level (Fig. 5e). In addition, both \textit{Kip3-CC} and 2x \textit{Kip3-Δ-T-LZ} localized to pre-anaphase and anaphase spindles similarly to \textit{Kip3} (Fig. 5f, 5g), consistent with a direct effect of these proteins on spindle length. These data support the conclusion that the sliding activity of \textit{Kip3} increases spindle length whereas the depolymerizing activity of \textit{Kip3} decreases it. These two activities counteract each other to regulate spindle length.

The sliding activity of \textit{Kip3} promotes spindle assembly in combination with kinesin-5/Cin8

In addition to its role in spindle length control, genetic analysis suggested that the microtubule sliding activity of \textit{Kip3}, in combination with that of Cin8/kinesin-5, makes an important contribution to bipolar spindle assembly. Individually, loss of Cin8 (\textit{cin8} \textit{Δ}) or the sliding activity of \textit{Kip3} (2x \textit{kip3} \textit{Δ-T-LZ}) did not compromise bipolar spindle assembly. However, \textit{cin8} \textit{Δ} 2x \textit{kip3} \textit{Δ-T-LZ} double mutants displayed a marked defect in bipolar spindle assembly (Fig. 6a, 6b). By contrast, a synergistic defect in bipolar spindle assembly was not observed by combining 2x \textit{kip3} \textit{Δ-T-LZ} with the deletion of the other kinesin-5 in budding yeast \textit{Kip1} (Fig. 6c, 6d). This is consistent with previous results suggesting \textit{Kip1} plays a less important role than Cin8 in spindle assembly\textsuperscript{36}. Additionally, we noted the appearance of diploids or near-diploid aneuploids in the population of \textit{cin8} \textit{Δ} 2X\textit{kip3} \textit{Δ-T-LZ} haploid strains (Fig. 6e). The increase in ploidy is likely explained by the defects in bipolar spindle assembly leading to a failure of karyokinesis. This effect was robust to alteration of \textit{kip3} \textit{Δ-T-LZ} gene dosage because diploids were also spontaneously generated in the population of \textit{cin8} \textit{Δ} \textit{kip3} \textit{Δ-T-LZ} haploid strains (Fig. 6e). Collectively these experiments establish that the sliding activity we report here for \textit{Kip3} makes physiologically important contributions to bipolar spindle assembly.

The two kinesin-5s, Cin8 and \textit{Kip1}, have been considered to be the major driving force for spindle assembly and spindle elongation in budding yeast. We therefore tested whether the sliding activity of \textit{Kip3} could replace the role of these two kinesin-5s. Bipolar spindles fail to assemble when both Cin8 and \textit{Kip1} are functionally inactivated\textsuperscript{36}. Overexpressing \textit{Kip3-CC} did partially rescue spindle collapse in cells lacking functional kinesin-5s (Fig. 6f). The fact that this rescue is modest is probably due to the optimization of kinesin-5s for spindle assembly, in terms of both their bipolar tetrameric structure and the characteristics of their motor domains, as previously proposed based on experiments using motor chimeras in the \textit{Xenopus} extract system\textsuperscript{37}.

Spindle elongation in budding yeast has an initial fast elongation phase (anaphase I) followed by a slow elongation phase (anaphase II). Cells lacking Cin8 have slower phase I rates whereas cells lacking \textit{Kip1} have slower phase II rates\textsuperscript{38}. Expression of \textit{Kip3-CC} from the native \textit{KIP3} promoter increased the phase I spindle elongation rate of \textit{cin8} \textit{Δ} cells as well as the phase II spindle elongation rate of \textit{kip1} \textit{Δ} cells (Fig. 6g). Thus, the sliding activity of \textit{Kip3}, if unopposed by its microtubule destabilizing activity, can partially substitute for individual kinesin-5s to promote spindle elongation.
DISCUSSION

We propose a “slide-disassemble” model to describe how Kip3 promotes both spindle assembly early in mitosis and spindle disassembly during mitotic exit (Fig. 7). At the beginning of mitosis, Kip3 contributes to bipolar spindle assembly through its anti-parallel microtubules sliding activity. This is supported by genetic data showing that the sliding activity of either Kip3 or Cin8 is required for bipolar spindle assembly. Measurements of spindle lengths in cells expressing Kip3 mutants that are selectively defective for either sliding or depolymerizing suggest that these activities balance in a way that Kip3 has no net effect on spindle length in pre-anaphase. However, in late anaphase the balance shifts, favoring microtubule destabilization and spindle disassembly. This “flip” from a positive role in spindle assembly to triggering spindle disassembly may be explained by changes in the amount of microtubule overlap and changes in microtubule length that occur during anaphase. As anaphase proceeds, the mass of overlapping microtubules decreases, and correspondingly, the number of Kip3 molecules involved in microtubule sliding decreases. At the same time, the length of individual microtubules becomes longer during anaphase, resulting in strong accumulation of Kip3 on microtubule plus ends, further shifting the balance towards microtubule destabilization. The net effect would be for the destabilizing activity of Kip3 to dominate at the end of anaphase, promoting spindle disassembly.

The microtubule sliding and cross-linking activities of Kip3 are likely to be relevant to the function of kinesin-8s in other eukaryotic cell types. RNAi experiments showed that the human kinesin-8s, Kif18A and Kif19, cooperate with kinesin-5/Eg5 to maintain spindle bipolarity in HeLa cells. This could result from an underlying mechanism similar to Kip3 because the tail domain of Kif18A binds microtubules in vitro and associates with microtubules in cells. In Drosophila spermatocytes, kinesin-8 Klp67A is likewise required to focus the central spindle during late anaphase. This is plausibly explained by a putative microtubule cross-linking activity of Klp67A because Klp67A also contains a secondary microtubule binding tail. We propose that the complex interplay between plus end directed motility, plus end destabilizing activity, microtubule cross-linking, and microtubule sliding will be a general feature of kinesin-8 family motors. Other microtubule regulators, such as XMAP215, dynein, and kinesin-14, may have similarly complex, context dependent functions (reviewed in [3]). An accurate understanding of mitotic spindle function will ultimately require an understanding of the complexity of its individual components.

METHODS

Strains and coding sequences for recombinant proteins

Supplementary Table S2 provides a list of yeast strains used in this study. Yeast strains were in the S288C background except those used for protein purification, which were in the W303-1A background. To fluorescently label the recombinant Kip3 variants, Halo tag (Promega), which enables covalently attachment of fluorescent dyes to proteins of interests in a 1:1 stoichiometry, was fused to the coding sequence of proteins through a GSSGSS linker.

Constructs and protein purification

The coding sequence for the N-terminal Halo-tagged full-length Kip3 (1-805), Kip3 ΔT –LZ (Kip3 1-480 + Gcn4 250-281) and Kip3-CC (Kip3 1-448 + Gcn4 253-281 + Kip3 478-805) were expressed from the GAL1 promoter on 2 μplasmids (pRS425 backbone). A protease-deficient yeast strain was used for protein expression. Protein expression was induced by addition of 2% galactose for 18h at 30°C prior to harvesting cells. Cells were disrupted by mechanical force using a coffee grinder. The yeast powder was thawed, and dissolved in...
cold lysis buffer containing 50 mM NaPO$_4$, 500 mM NaCl, 10% glycerol, 0.5 mM ATP-Mg, 40 mM imidazole, 2 mM DTT, 1% Triton X-100, protease inhibitor cocktail tablets (Roche), and 2mM PMSF at pH 7.4. Dounce homogenization was used to further lyse cells at 4°C. After centrifugation at 45,000 rpm for 45 min at 4°C, the supernatant was collected and incubated with Ni$^{2+}$ sepharose (GE Healthcare) for 30 min at 4°C. The beads were washed 3 times with a buffer containing 50 mM NaPO$_4$, 500 mM NaCl, 10% glycerol, 0.1 mM ATP-Mg, 25 mM imidazole, 0.1 mM DTT, protease inhibitor cocktail tablets, and 1 mM PMSF at pH 7.4. The protein was labeled with 10 μM TMR (Promega) on beads for 1.5 h at 4°C before elution of the beads with stripping buffer (50 mM HEPES, 300 mM NaCl, 50 mM EDTA, 5 mM MgCl$_2$, 50 μM ATP-Mg, and 0.1 mM DTT at pH 7.4). The protein elutes were adjusted to 250 mM NaCl and loaded onto a Uno S1 (Bio-rad) ion exchange column, which had been equilibrated with 50 mM HEPES, 250 mM NaCl, and 50 μM ATP-Mg at pH 7.4. The protein was eluted by a linear gradient to 1M NaCl over 15 column volumes. The elutes were supplemented with 10% sucrose and centrifuged at 80,000 rpm for 5 min at 4°C. The supernatant was collected and snap-frozen in liquid nitrogen and stored at -80°C.

**Chemical cross-linking**

Proteins around 1 μM were incubated with freshly prepared 1 mM ethylene glycol-bis(succinic acid N-hydroxysuccinimide ester) (EGS, Sigma) at room temperature for 30 min in 50 mM HEPES, 400 mM NaCl, 50 μM ATP-Mg, 10% sucrose at pH 7.4. The reactions were quenched by adding 40 mM Tris-Cl, 50 mM glycine at pH 7.5 and incubating at room temperature for 30 min.

**Preparing stabilized microtubules**

Tubulin was purified from bovine brain by two rounds of assembly and disassembly, followed by phosphocellulose chromatography. NEM (N-ethylmaleimide) modified tubulin was prepared using the Mitchison Lab protocol (http://mitchison.med.harvard.edu/protocols/polarity.html). Fluorescein, rhodamine, HiLyte 488, HiLyte 647, and biotin -labeled tubulin were purchased from Cytoskeleton. To prepare Taxol-stabilized microtubules for motility assays, a tubulin stock [BRB80 (80 mM PIPES-K$^+$, 1 mM EGTA, 1 mM MgCl$_2$ at pH 6.8) containing 20 μM tubulin, 2 μM fluorescein-labeled tubulin, 2 μM biotin-labeled tubulin, 1 mM GTP-Mg, 1 mM DTT, and 10% DMSO] was incubated at 37°C for 45 min. Taxol was added to a concentration of 20 μM. The mix was then incubated at 37°C for 3 hr prior to use. To prepare GMPCPP-stabilized microtubules for the depolymerase assay, microtubule seeds were generated by incubating the seed mix (BRB80 containing 20 μM tubulin, 0.2 μM biotin-tubulin, 5 μM fluorescein-tubulin, and 1 mM GMPCPP-Mg) at 37°C for 30 min. The elongation mix was made by diluting the seed mix at 1:10 ratio with BRB80+0.5 mM GMPCPP. The seed was added to the elongation mix at a 1:40 ratio after pre-warming the elongation mix at 37°C for 1 min. The final mix was incubated for 3 h at 37°C before use. To prepare long track microtubules for the sliding assays, a tubulin seed stock [BRB80 (80 mM PIPES-K$^+$, 1 mM EGTA, 1 mM MgCl$_2$ at pH 6.8) containing 20 μM tubulin, 2 μM fluorescein-labeled tubulin, 2 μM biotin-labeled tubulin, and 1 mM GTP-Mg] was thawed, supplemented with 10% DMSO, and incubated at 37°C for 45 min. Taxol was added to a concentration of 20 μM. The mix was then incubated at 37°C for 3 hr or longer. An elongation mix stock [BRB80 containing 2 μM tubulin, 0.5 μM fluorescein-labeled tubulin, 0.02 μM biotin-labeled tubulin, 0.5 μM NEM-tubulin, and 1 mM GMPCPP-Mg] was pre-warmed to 37°C, and then added into the seed mix for 1h incubation at 37°C before ready to use. A similar procedure was adopted to prepare the short cargo microtubules, with the following modifications: 1. Fluorescein tubulin was replaced with rhodamine tubulin or HyLite 647 tubulin; 2. Biotin-tubulin was omitted. 3. GMPCPP was used in all reactions. 4. The incubation time for the seed mix was shortened to 30 min.
**Sliding assay on stabilized microtubules**

Assay chambers with a volume of approximately 10 μL were made by forming a “coverslip-double-sided tape-glass slide” sandwich. The buffer volume flowed into the chamber in each of the following steps are around 20 μL. The chambers were sequentially coated with 1 mg/mL biotin-BSA and 0.5 mg/mL streptavidin, allowing Taxol-stabilized fluorescein-labeled track microtubules to be immobilized on the coverslips. The reaction pre-mixtures contained BRB80 buffer at pH 7.2 supplemented with 0.5 mg/mL casein, 5% glycerol, 20 μM Taxol, 50 mM KCl. Pre-mixtures supplemented with 0.5 mM AMPPNP and 150 nM motors were flowed into the chamber, and then rhodamine-labeled cargo microtubules diluted in the pre-mixtures supplemented with 0.5 mM AMP-PNP were flowed into the chambers. AMPPNP was used to enhance the frequency that cargo microtubules were captured by motors on track microtubules. Microtubule alignment and sliding were also observed with ATP but at a lower frequency. After cargo microtubules were captured by motors in the chamber, the chamber was washed with pre-mixtures supplemented with 0.5 mM AMP-PNP. The final reaction buffer contained the pre-mixtures supplemented with 2 mM ATP and oxygen scavenging mix (0.2 mg/mL glucose oxidase, 0.035 mg/mL catalase, 25 mM glucose and 70 mM β-mercaptoethanol). An image from the GFP channel was taken at the beginning of the experiment to locate the position of the track microtubules. Time-lapse images from the Cy3 channel were acquired every 15 s for 20 min to trace the movement of cargo microtubules. The images were captured using a Zeiss AX10 Imager M1 microscope equipped with a CoolSnap HQ camera (Roper Scientific) and a 63× 1.4 N.A. oil immersion objective. The same microscope, camera, and objective were used for all the other imaging experiments except those involving TIRF microscopy. All the images were captured at room temperature unless specified. Kymographs were generated to quantify the sliding speeds, motor velocity, run length, end dwell time, and depolymerase rates.

**Sliding of dynamic microtubules growing from micropatterned seeds**

Detailed methods were previously described28. This protocol was followed except that the composition of the elongation mix was modified: 2 μM of Atto-488 tubulin, 8 μM of unlabeled tubulin in BRB80 supplemented with 50 mM NaCl, 1 mM GTP, 0.2% BSA, 0.025% methyl cellulose (1500 CP), and an oxygen scavenger cocktail (2 mg/mL glucose, 80 μg/mL catalase, and 0.67 mg/mL glucose oxidase). Dynamic microtubules were visualized using a 60x objective-based azimuthal ilas2 TIRF microscope (Nikon eclipse Ti, modified by Roper scientific) equipped with an Evolve 512 camera (Photometrics). The microscope stage was maintained at 32°C using a warm stage controller (LINKAM MC60). Excitation was achieved using 491 nm and 561 nm lasers (Optical Insights) with an exposure time of 70 ms. Time-lapse recording (one frame every 5 s) was performed using Metamorph® software (version 7.7.5, Universal Imaging). Movies were processed to improve signal/noise ratio (equalize light, low pass and flatten background filters of Metamorph® software). Static images were taken 30 min after the reactions were started in order to quantify the microtubule bundle deflections.

**Motility assays**

Track microtubules were immobilized in the assay chamber as described above. Motor proteins were diluted to 50 pM in reaction mixtures containing BRB80 buffer at pH 7.2 supplemented with 0.5 mg/mL casein, 5% glycerol, 2 mM ATP-Mg, 20 μM Taxol, 50 mM KCl, and oxygen scavenging mix (0.2 mg/mL glucose oxidase, 0.035 mg/mL catalase, 25 mM glucose and 70 mM β-mercaptoethanol). Images were recorded with 100 ms exposure every 2 s for 20 min, using an Olympus IX-81 TIRF microscope equipped with a back-thinned electron multiplier CCD camera (Hamamatsu) and a 100× 1.45 N.A. oil immersion TIRF objective.
Depolymerase assay

Experimental conditions were similar to those for the motility assay, with the following modifications: GMPCPP-stabilized microtubules were used instead of Taxol-stabilized microtubules. Taxol was not included in the reaction mixtures. Wide-field images were acquired every 15 s for 10 min. Motor input was from 6 nM to 24 nM.

Dynamic microtubules assay

Flow cells were made using double-sided tape with passivated coverslips containing PEG and PEG-Biotin (2 mg/ml and 20 μg/ml respectively, Laysan Bio) as described in 47. HiLyte 488-labeled GMPCPP seeds were immobilized to the coverslips as described in the motility assay. The tubulin growth mix (14 μM tubulin, 0.7 μM HiLyte 647-labeled tubulin, 1 mM GTP, 2 mM ATP-Mg, 1 mg/ml casein, 0.1% methyl cellulose and an oxygen scavenging system described in the motility assay) was then introduced into the chamber. Microtubules were allowed to grow for 5 min at room temperature prior to flowing in motors diluted in the same growth mix. Time-lapse imaging was recorded at room temperature with 100 ms exposure every 0.5 s for 10 min. Image analysis was performed as previously described in 14. Pauses were included in the calculation of growth rates.

Cellular imaging and data analysis

Yeast strains were grown to log phase in SC media at 30°C prior to imaging. To measure the spindle length in cells overexpressing Kip3 variants, cells were arrested in S phase by treating with 0.1 mM hydroxyurea for 3 hours, followed by replacing 2% Raffinose with 2% Galactose in the media, and incubating for another 2.5 hours in the presence of hydroxyurea before imaging. To determine the timing of bipolar spindle formation, cells were synchronized by treatment of 20 μM alpha factor for two hours. Cells were then released from the G1 block and collected at the indicated timepoints. 3.7% paraformaldehyde was then added for 8 min at room temperature, washed with PBS, and stored at 4°C prior to imaging. To determine if overexpressing Kip3-CC rescues the defects in forming bipolar spindles in cin8-3 kip1 Δ cells, cells were grown to log phase in SC media supplemented with 2% raffinose at 24°C and then arrested in S phase by 0.1 mM hydroxyurea for 3 h. The expression of Kip3 variants were induced by replacing 2% raffinose with 2% galactose for 3 h. Finally cells were shifted to 37°C for 1 h to inactivate Cin8 before imaging. To perform time-lapse imaging to monitor astral microtubule dynamics and spindle elongation dynamics, a “coverslip - double-sided tape - glass slide” sandwich chamber of 50 μL was used and pre-coated with 0.5 mg/mL Concanavalin A. Cells were then flowed into the chamber and the chamber was inverted to let cells immobilized to the coverslip. Unbounded cells were then washed away by three volumes of SC media prior to imaging.

Cellular images were acquired using a Zeiss AX10 microscope equipped with a CoolSnap HQ camera (Roper Scientific). A Z-focal plane series of images (7 planes with 0.75 μm between adjacent planes) was captured with a 63× 1.4 N.A. oil immersion objective. The spindle lengths were determined by measuring the distance between the centroid of Spc42-marked spindle pole bodies in 3D. The fluorescence intensity of individual proteins was measured from a maximal intensity projection from the Z-stack. Cellular background was subtracted for quantification. To monitor astral microtubule dynamics, images were acquired every 6 s for 10 min. Microtubule growth rates and shrinkage rates were obtained by fitting the time-length plot to a linear regression line. Microtubule catastrophe and rescue rates were calculated by dividing the number of events by the total microtubule life time. To monitor spindle elongation dynamics, images were acquired every 1 min for 90 min. Spindle elongation rates were obtained by fitting the time-length plot to a linear regression line.
Western blotting

Standard procedures were used for Western blotting. Mouse anti-GFP antibody (7.1 and 13.1, Roche, 11814460001) was used in a 1:1,000 dilution. Rat anti-α-tubulin antibody (YOL 1/34, Accurate Chemical & Scientific, YSRTMCA78G) was used in a 1:3,000 dilution.

Flow cytometry

Cells were grown to log phase and fixed in 70% ethanol for two hours. Cellular RNA was digested by 0.25 mg/mL RNase in 50 mM sodium citrate, pH 7 overnight at 37°C. Then DNA was stained by 0.015 μg/μL propidium iodide for one hour at room temperature. FACS analysis was performed with a FACScalibur flow cytometer (Becton Dickinson).

Statistical analysis

Significance was calculated using a two-tailed Student’s t-test. A P value of less than 0.05 was considered statistically significant. Values are reported as mean ± s.e.m.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Kip3-mediated microtubule sliding activity

a) Schematic of the constructs used in this study. Kip3 contains a head (red, also called motor domain), followed by a short neck consisting of 4-5 coiled-coil heptads (blue), and a tail domain (orange). The leucine zipper (green) was added to maintain the dimeric state of the tail-less Kip3 (Kip3ΔT-LZ)\textsuperscript{21}. Kip3-CC was made by replacing the native Kip3 neck with the leucine zipper.

b) Cargo microtubules (red, HiLyte 647 labeled) were aligned with track microtubules (green, HiLyte 488 labeled) in the presence of Kip3, but not in the presence of Kip3ΔT-LZ. Track microtubules were first immobilized on the coverslip. Kip3 or Kip3ΔT-LZ at 150 nM, cargo microtubules, and a final reaction mix were sequentially flowed into the chamber. 20 μM Taxol was included in the assay to prevent microtubule depolymerization by Kip3. In this assay, a wash was additionally applied after motors’ binding to the track microtubules, reducing non-specific adherence of motors to coverslips. Scale bar: 5 μm.
c) Kymographs show the localization of Kip3 during sliding or “tug-of-war” movements. Kip3 was labeled with tetramethylrhodamine and cargo microtubules were labeled with HiLyte 647. Input: 150 nM Kip3. Scale bar: 2 μm (horizontal) and 2 min (vertical).

d) Microtubules in a parallel orientation display “tug-of-war” movement in the presence of 150 nM of Kip3. Polarity-labeled microtubules (bright red for the plus end of cargo microtubules and bright green for the plus end of track microtubules) show parallel orientation. “+” and “-” marked the microtubule plus end and minus end, respectively. Scale bar: 5 μm. Kymographs at the bottom show “tug-of-war” movement of cargo microtubules with the numbers corresponding to the number in the images above. Scale bar: 2 μm (horizontal) and 2 min (vertical).

e) Anti-parallel microtubule sliding by Kip3. Same figure settings and experimental conditions as in d).

f) Kip3 slides cargo microtubules (arrows) along track microtubules with a similar velocity to that with which Kip3 glides cargo microtubules (arrow head). (LEFT) The majority of cargo microtubules (red) were aligned with track microtubules (green) in the presence of 150 nM Kip3. Unaligned cargo microtubules were occasionally observed, which was captured by those Kip3s that non-specifically attached to coverslips. Scale bar: 5 μm. (RIGHT) Quantification of microtubule velocity with an initial input of 150 nM Kip3. Shown are mean ± s.e.m. (N=50 microtubules).
Figure 2. Kip3-mediated sliding of dynamic microtubules

a) Schematic of the method for micropatterned microtubule growth. Polyethylene glycol (PEG) - coated glass was placed under a photomask. The exposed spots, spaced by 20 μm, were made competent for the adsorption of microtubule seeds by deep UV induced oxidation of PEG. Tubulin was added to allow polymerization of dynamic microtubules from the seeds.

b) Time-lapse imaging showing curved microtubule bundles induced by Kip3. Dynamic microtubules (green) were growing from seeds (red) adsorbed on micropatterned sites in the presence of 10 μM of tubulin together with the indicated proteins (50 nM of Kip3, 33 nM of Ase1). White arrows indicate curved microtubule bundles in the presence of Kip3. Scale bar: 10 μm.

c) Quantification of the microtubule bundle deflection as an indicator of microtubule sliding. (LEFT) Schematic of the quantification. (RIGHT) Kip3 significantly increases the deflection of bundles. (N=41, 40, 190, 324 microtubule bundles for control, Kip3, Ase1, and Kip3+Ase1, respectively).

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Figure 3. Kip3-CC is a sliding proficient but depolymerizing deficient Kip3 variant

a) Kip3-CC slides microtubules. (TOP) Cargo microtubules (red) were aligned with track microtubules (green) in the presence of Kip3-CC. (BOTTOM) Shown is a Kymograph of a single track microtubule with one cargo showing a “tug-of-war” movement (left) and another cargo exhibiting sliding movement (right). Kip3-CC at 150 nM, cargo microtubules, and a reaction mix were sequentially added. Scale bar: 5 \( \mu \)m (horizontal) and 2 min (vertical).

b) Kymographs show the localization of Kip3-CC during sliding or “tug-of-war” movements. Kip3-CC was labeled with tetramethylrhodamine and cargo microtubules were labeled with HiLyte 647. Input: 150 nM Kip3-CC. Scale bar: 2 \( \mu \)m (horizontal) and 2 min (vertical).

c) Quantification of sliding rates with an initial input of 150 nM Kip3 or Kip3-CC. Shown are mean ± s.e.m. (N=50 microtubules).

d) Kip3-CC is deficient for depolymerase activity. (TOP) Kymographs from depolymerase assays using GMPCPP-stabilized microtubules as substrates. Protein input: 12 nM. (BOTTOM) Quantification of the microtubule depolymerization rates of Kip3 or Kip3-CC.
Because Kip3 depolymerizes microtubules in a length-dependent manner, microtubules within a narrow range of lengths were chosen for quantification. 3-5 μm were chosen to correspond to the range of microtubule lengths in budding yeast (mostly below 5 μm). Shown are mean ± s.e.m. (N=50 microtubules). Scale bar: 2 μm (horizontal) and 1 min (vertical).

e) Kip3-CC is defective in promoting catastrophe of dynamic microtubules in vitro. (TOP) Kymographs show the assembly of HiLyte-647 labeled dynamic microtubules. Because Kip3-CC has higher microtubule landing frequency than Kip3 in the presence of tubulin dimers, the protein input was adjusted (150 nM Kip3 Vs 6 nM Kip3-CC) to achieve a nearly identical measured flux of Kip3 and Kip3-CC on microtubules. Scale bar: 5 μm (horizontal) and 1 min (vertical). (BOTTOM) Quantification of the catastrophe frequency in the assay described above. Shown are mean ± error. (N=164, 25, and 23 catastrophe events for Kip3, Kip3-CC, and control, respectively). Growth rate and shrinkage rate are shown in Supplementary Fig. S7c.
Figure 4. Kip3-CC retains comparable motility to Kip3

a) Kymographs from single molecule imaging assay visualizing the motility of TMR-labeled Kip3 or Kip3-CC at ~ 0.1 nM on Taxol-stabilized microtubules. An image of fluorescein-labeled microtubules was acquired separately to determine the position of microtubule ends, as indicated by asterisks. Scale bar: 5 μm (horizontal) and 1 min (vertical).

b) Quantification of the motility and plus end dwell time of Kip3 and Kip3-CC. Histograms show the distribution of motors’ velocity, run length, and plus end dwell time. The distribution of velocity was fit to a Gaussian curve. The distribution of run length and plus end dwell time were fit to a first-order exponential curve. Shown are mean ± s.e.m. (N=150 – 250 motors).
Figure 5. Counterbalancing effects between the sliding and destabilizing activity of Kip3 in the control of spindle length

a) Fluorescent microscopy visualizing spindles in cycling cells overexpressing Kip3-CC. Green: Tub1-GFP marked spindle, Red: Spc42-CFP marked spindle pole body, Blue: cell boundary obtained by bright field images. Kip3-CC was overexpressed using a pGAL1 expression system. Scale bar: 5 μm.

b) Comparisons of spindle lengths and astral microtubule lengths among cells overexpressing the indicated constructs. These cells were synchronized in S phase by hydroxyurea. Shown are mean ± s.e.m. (N=50 cells).

c) Comparison of pre-anaphase spindle lengths in cells expressing kip3 mutants from the endogenous KIP3 promoter. Tub1-GFP was used as a tracer to visualize spindles. Shown are mean ± s.e.m. (N=50 cells). P values were obtained from a two-tailed Student’s t-test.
d) Effects of Kip3 mutants on late anaphase spindle length. Time-lapse imaging was performed to measure spindle length just prior to spindle disassembly. To exclude the effect from cell size on spindle length, we measured cell length on the long axis and verified that there were no significant differences among the size of four strains. Shown are mean ± s.e.m. (N=15 cells). P values were obtained from a two-tailed Student’s t-test.

e) Steady-state protein levels of EYFP-tagged Kip3 and variants. Shown is a western blot using an anti-GFP antibody. Tubulin serves as a loading control.

f) Localization of Kip3 and variants throughout the cell cycle. Images from fluorescence microscopy show cells expressing one copy of Kip3, one copy of Kip3-CC and two copies of Kip3 ΔT-LZ. Kip3 and variants were labeled with EYFP. Microtubules were labeled with CFP-Tub1. Images of single channel of Kip3 and variants were shown in the top panel. Merged images were shown in the bottom panel. Scale bar: 2 μm.

g) Quantification of accumulation of Kip3 and variants to pre-anaphase spindles. Shown are mean ± s.e.m. (N=50 spindles)
Figure 6. The sliding activity of Kip3 promotes spindle assembly

a) Bipolar spindles were formed in large-budded cells from wild-type strains whereas the dot-like monopolar spindles were found in large-budded cells from 2x kip3ΔT-LZ cin8Δ strains. Spindles are labeled with GFP-Tub1. Scale bar: 5 μm.

b) Synergistic defects between 2x kip3ΔT-LZ and cin8Δ in forming bipolar spindles. Cells were arrested in the G1 phase by α-factor treatment and then released from the arrest. The percentage of cells with bipolar spindles was measured at the indicated time points after release. Two hundred cells were scored for each data point.

c) Images showing bipolar spindles formed in wild-type and 2x kip3ΔT-LZ kip1Δ strains. Spindles are labeled with GFP-Tub1. Scale bar: 5 μm.

d) 2x kip3ΔT-LZ and kip1Δ cells have a subtle defect in bipolar spindle assembly. Cells were arrested in the G1 phase by α-factor treatment, released, and the percentage of cells
with bipolar spindles was quantified at the indicated time points. Two hundred cells were scored for each data point.
e) Significant increase in the ploidy of cells containing \textit{kip3} \textit{Δ} T-LZ \textit{cin8} \textit{Δ} double mutants. DNA content of the indicated strains was monitored by FACS sorting of propidium iodide stained cells. Note the 4N peak (arrows) appearing in 2x \textit{kip3} \textit{Δ} T-LZ \textit{cin8} \textit{Δ} cells and \textit{kip3} \textit{Δ} T-LZ \textit{cin8} \textit{Δ} cells, which indicates the presence of diploids in the population.
f) Overexpression of Kip3-CC partially rescues the spindle collapse of strains compromised for kinesin-5 function. \textit{cin8-3 kip1} \textit{Δ} cells were arrested in S phase by hydroxyurea at 24°C. The expression of Kip3 variants was then induced. Aliquots of cells were either maintained at 24°C or shifted to 37°C to inactivate Cin8. Spc42-CFP and Tub1-GFP enabled the visualization of spindle pole bodies and spindles. Two hundred cells were scored for each condition. N= 3 independent experiments. Shown are mean ± s.e.m. P values were obtained from a two-tailed Student’s t-test.
g) The sliding activity of Kip3 promotes spindle elongation in anaphase. Kip3-CC was expressed under endogenous \textit{KIP3} promoter. Time-lapse imaging was used to monitor spindle elongation in anaphase, which can be split into a fast phase (phase I) and a slow phase (phase II). Shown is mean ± s.e.m. (N=15 cells). #: P>0.05, *: 0.005<P<0.05, **: P<0.005. P values were obtained from a two-tailed Student’s t-test.
Figure 7. A ‘slide-disassemble’ model to explain the complex effects of Kip3 on regulating spindle length and stability

(TOP) Cartoon illustrates how the microtubule sliding and destabilizing activity of Kip3 may affect spindle length individually and in combination. In wild-type pre-anaphase cells, sliding and destabilizing activities appear to be in approximate balance, a conclusion that is supported by the absence of an alteration in pre-anaphase spindle length in cells completely lacking Kip3. Mutations that tip the balance in one direction or the other have corresponding effects on spindle length (long spindles in kip3-CC cells and shorter spindles in 2x kip3ΔT-LZ cells). Moreover, when the balance is tipped toward the destabilizing slide, as in 2x kip3ΔT-LZ cells, bipolar spindles fail to assemble if microtubule sliding is further diminished by loss of another anti-parallel sliding motor Cin8/kinesin-5.

(BOTTOM) Upon anaphase entry, the numbers of overlapping microtubules decreases and the degree of overlap for microtubule pairs eventually shrinks. At the same time, the length of interpolar microtubule increases by about five-fold, which is predicted to increase the
amount of plus end-associated Kip3 because of Kip3’s remarkable processivity. The increased amount of Kip3 will destabilize interpolar microtubules and eventually promote spindle disassembly.