

Five factors can reconstitute all three phases of microtubule polymerization dynamics

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Cytoplasmic microtubules (MTs) undergo growth, shrinkage, and pausing. However, how MT polymerization cycles are produced and spatiotemporally regulated at a molecular level is unclear, as the entire cycle has not been recapitulated *in vitro* with defined components. In this study, we reconstituted dynamic MT plus end behavior involving all three phases by mixing tubulin with five *Drosophila melanogaster* proteins (EB1, XMAP215^{Msps}, Sentin, kinesin-13^{Klp10A}, and CLASP^{Mast/Orbit}). When singly mixed with tubulin, CLASP^{Mast/Orbit} strongly inhibited MT catastrophe and reduced the growth rate. However, in the presence of the other four factors, CLASP^{Mast/Orbit} acted as an inducer of pausing. The mitotic kinase Plk1^{Polo} modulated the activity of CLASP^{Mast/Orbit} and kinesin-13^{Klp10A} and increased the dynamic instability of MTs, reminiscent of mitotic cells. These results suggest that five conserved proteins constitute the core factors for creating dynamic MTs in cells and that Plk1-dependent phosphorylation is a crucial event for switching from the interphase to mitotic mode.

Introduction

Complementary to genetic analysis, *in vitro* reconstitution of intracellular processes, using defined factors, is a prerequisite to understand the process at a molecular level. This can determine if a given effect is directly mediated by each factor and defines a minimal set of factors that can execute the process (Loisel et al., 1999; Garner et al., 2007; Shintomi et al., 2015; Yeeles et al., 2015). Dynamic microtubules (MTs) are composed of tubulin proteins and play critical roles in various intracellular processes in eukaryotic cells, such as spindle formation and cell polarization. Tubulin proteins, at particular concentrations, can autonomously and stochastically polymerize and depolymerize *in vitro*; this is referred to as MT dynamic instability (Mitchison and Kirschner, 1984; Horio and Hotani, 1986; Walker et al., 1988). In cells, free MT plus ends in the interphase cytoplasm repeatedly undergo three phases (growth, shrinkage, and pausing) in a stochastic order (Shelden and Wadsworth, 1993; Rogers et al., 2002). Multiple proteins control cellular MT dynamics, and depletion of these proteins results in alterations to MT dynamicity and consequently a change in the length of MT-based higher-order structures such as cilia or the mitotic spindle (Howard and Hyman, 2007; Akhmanova and Steinmetz, 2008; Goshima and Scholey, 2010; Hu et al., 2015). Most, if not all, of these proteins can individually modulate MT dynamics *in vitro*. Examples include XMAP215, which can increase MT growth rate, or kinesin-13, which can increase “catastrophe” frequency (frequency of growth-to-shrinkage or pause-to-shrinkage transitions; Howard and Hyman, 2007). When two or more factors are mixed with tubulin, the dynamic MT behavior is better

reproduced (Kinoshita et al., 2001; Li et al., 2012; Zanic et al., 2013). However, cellular MT dynamics, with all three phases, have not been successfully reproduced *in vitro*. Moreover, upon entry into mitosis, MTs become more dynamic, wherein MT growth rate and catastrophe frequency are increased and pausing is suppressed. However, the molecular basis of this cell cycle-dependent switch is also unclear.

The *Drosophila melanogaster* S2 cell line is a promising model system for the reconstitution of MT plus end dynamics. Parameters of MT dynamics in interphase and mitosis have been obtained through live-cell microscopy of fluorescent markers (Rogers et al., 2002; Brittle and Ohkura, 2005; Sousa et al., 2007; Li et al., 2011; Trogdon and Rogers, 2015). Using this cell line, a handful of proteins required for dynamic regulation of MTs have been identified through multiple functional studies, including a large-scale functional genomics screen (Goshima et al., 2007; Hughes et al., 2008; Moutinho-Pereira et al., 2013). For instance, cells depleted of three conserved MT plus end tracking proteins Msps (XMAP215/ch-TOG orthologue), EB1 (a member of the EB protein family), and Sentin (a functional homolog of mammalian SLAIN2) similarly decreased MT dynamicity, wherein MT growth rate was reduced and pausing was drastically elevated (Rogers et al., 2002; Brittle and Ohkura, 2005; Li et al., 2011). To initiate *in vitro* reconstitution of MT plus end dynamics in S2 cells, we purified EB1, Sentin, and XMAP215^{Msps} proteins, mixed them with pig brain-derived tubulins, and assessed MT plus end polymerization dynamics (Li

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Abbreviations used: dsRNA, double-stranded RNA; KD, kinase dead; MT, microtubule.

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et al., 2012). Compared with MTs generated solely with tubulin, or with one of these proteins, MTs formed in the presence of all three proteins had faster growth and more frequent catastrophe events, which is consistent with their depletion phenotype in cells (Rogers et al., 2002; Brittle and Ohkura, 2005; Li et al., 2011, 2012). However, even with various concentrations and combinations of these proteins, an in vitro system that closely reproduces the in vivo dynamics has not yet been constructed; most critically, pausing and the “rescue” event (shrinkage-to-growth or shrink-to-pause transitions) were observed with much lower frequency than in cells (Li et al., 2012).

We reasoned that a failure to reconstitute these three phases realistically might be caused by missing factors in the in vitro reaction. In this study, we considered Klp10A (kinesin-13) and CLASP (*Drosophila* name is Mast or Orbit) as likely candidates for these missing factors, because RNAi depletion in S2 cells affected MT dynamics and mitotic spindle size (Goshima et al., 2005; Mennella et al., 2005; Sousa et al., 2007). Furthermore, orthologous proteins of other species were shown to modulate MT dynamics in vitro (Mennella et al., 2005; Al-Bassam et al., 2010). In addition, with recently established methodology (Widlund et al., 2012), we purified tubulin proteins from S2 cells and used them to replace pig tubulin proteins. In the reaction, in vivo MT dynamics were reproduced qualitatively (and, to some extent, quantitatively) when five factors were mixed with S2 tubulin. Furthermore, phosphorylation by the mitotic kinase Polo (Plk1 in mammals) shifted MT dynamics to the mitotic mode.

Results

CLASP^{Mast/Orbit} suppresses MT growth and catastrophe

We first characterized the activity of S2 tubulin in the MT plus end polymerization assay, in which dynamic MTs are generated from a stable MT seed made with GMPCPP-tubulin (Bieling et al., 2007; Fig. 1 A). As expected, S2-derived tubulins showed repeated growth-shrinkage cycles, as observed with pig tubulins, although each parameter was quantitatively different (Fig. S1, A, C, and D).

Full-length kinesin-13^{Klp10A} was purified from *Escherichia coli*, and it showed potent MT depolymerization activity toward both ends of seed MTs (Fig. S1 E), as expected (Rogers et al., 2004; Mennella et al., 2005).

GFP-tagged CLASP^{Mast/Orbit} was purified from insect Sf21 cells (Fig. S1 A). Gel filtration chromatography indicated that the protein does not form aggregates (Fig. S1 B). When mixed with S2 tubulin and MT seeds, GFP-CLASP^{Mast/Orbit} was localized along the seed and dynamic MTs, and a dramatic reduction in MT growth rate and catastrophe frequency was observed (Fig. 1, B–D). Such activity was not observed, and MT association was reduced, when we used a mutant GFP-CLASP^{Mast/Orbit} in which four amino acid residues critical for MT or tubulin binding in the TOG2 and TOG3 domains were mutated (Al-Bassam et al., 2010; Al-Bassam and Chang, 2011; Leano et al., 2013; Fig. 1, B–D). Thus, the observed growth- and catastrophe-suppression activities were derived from CLASP^{Mast/Orbit}, and not from contaminated proteins of Sf21 cells. In addition, the result indicated that the effect of GFP-CLASP^{Mast/Orbit} (on MT growth rate and catastrophe) requires its binding to tubulin and/or MTs. Previous RNAi experiments suggested that CLASP^{Mast/Orbit}

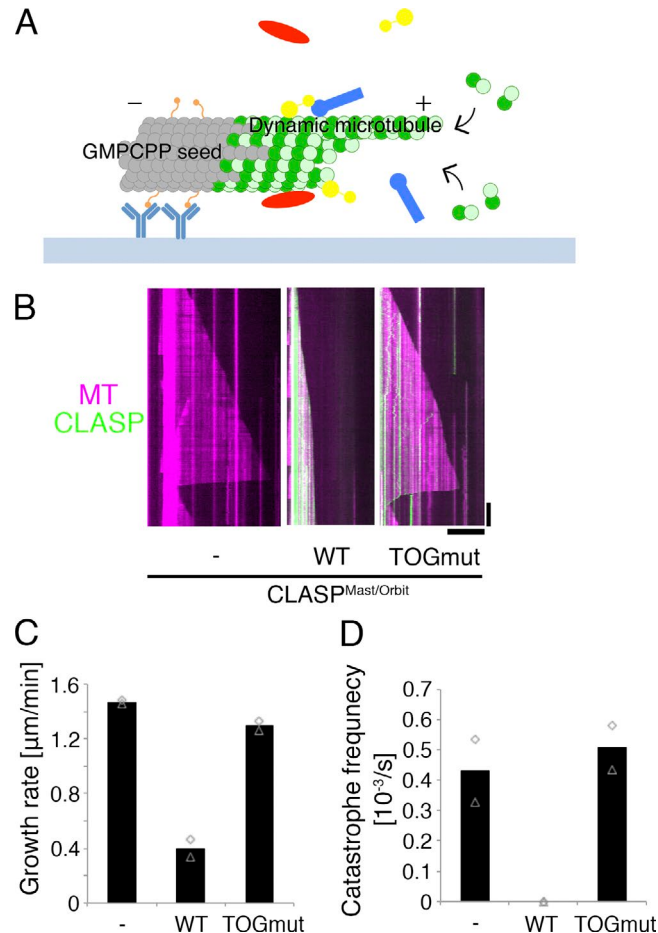


Figure 1. CLASP^{Mast/Orbit} slows growth and inhibits catastrophe in vitro. (A) Schematic of in vitro MT polymerization assay used in this study. Green, tubulin; orange, biotinyl; light blue, anti-biotin antibody; red/yellow/blue, recombinant proteins. (B) CLASP^{Mast/Orbit} slows MT growth and inhibits catastrophe. Kymographs show GFP-CLASP^{Mast/Orbit} (wild type [WT] and TOG domain mutant [TOGmut]) and MT (10 μ M S2 tubulin). (C and D) MT growth rate and catastrophe frequency in the presence of 45 nM CLASP^{Mast/Orbit} (wild type and TOG domain mutant) and 10 μ M S2 tubulin. The mean values of each experiment are marked in gray, whereas the mean values of all the experiments are indicated by black columns (two independent experiments). Actual values are plotted in these graphs ($n = 15\text{--}22$ MTs). Bars: (horizontal) 10 μ m; (vertical) 2 min.

induces MT pausing (Sousa et al., 2007; Trogdon and Rogers, 2015), whereas purified fission yeast CLASP^{Clsl} protein was shown to have potent rescue (shrinkage-to-growth) activity (Al-Bassam et al., 2010). However, we could not assess if *Drosophila* CLASP^{Mast/Orbit} has these activities in this experiment, because MT shrinkage was never observed (Fig. 1 D).

CLASP has an EB1-binding SxIP motif and abundantly localizes at the growing end of MTs in cells (Sousa et al., 2007; Honnappa et al., 2009). We reconstituted the plus end-tracking behavior of CLASP^{Mast/Orbit} by adding EB1 to the reaction (Fig. S1 F). EB1 on its own slightly elevated MT growth rate and catastrophe frequency (Li et al., 2012). However, we could not detect any additional effects of CLASP^{Mast/Orbit} on growth rate; in addition, catastrophe could not be observed regardless of the presence or absence of EB1 (Fig. S1 G). These results suggest that EB1-dependent plus end enrichment of CLASP^{Mast/Orbit} is not a prerequisite for the two CLASP^{Mast/Orbit}-mediated activities toward plus ends.

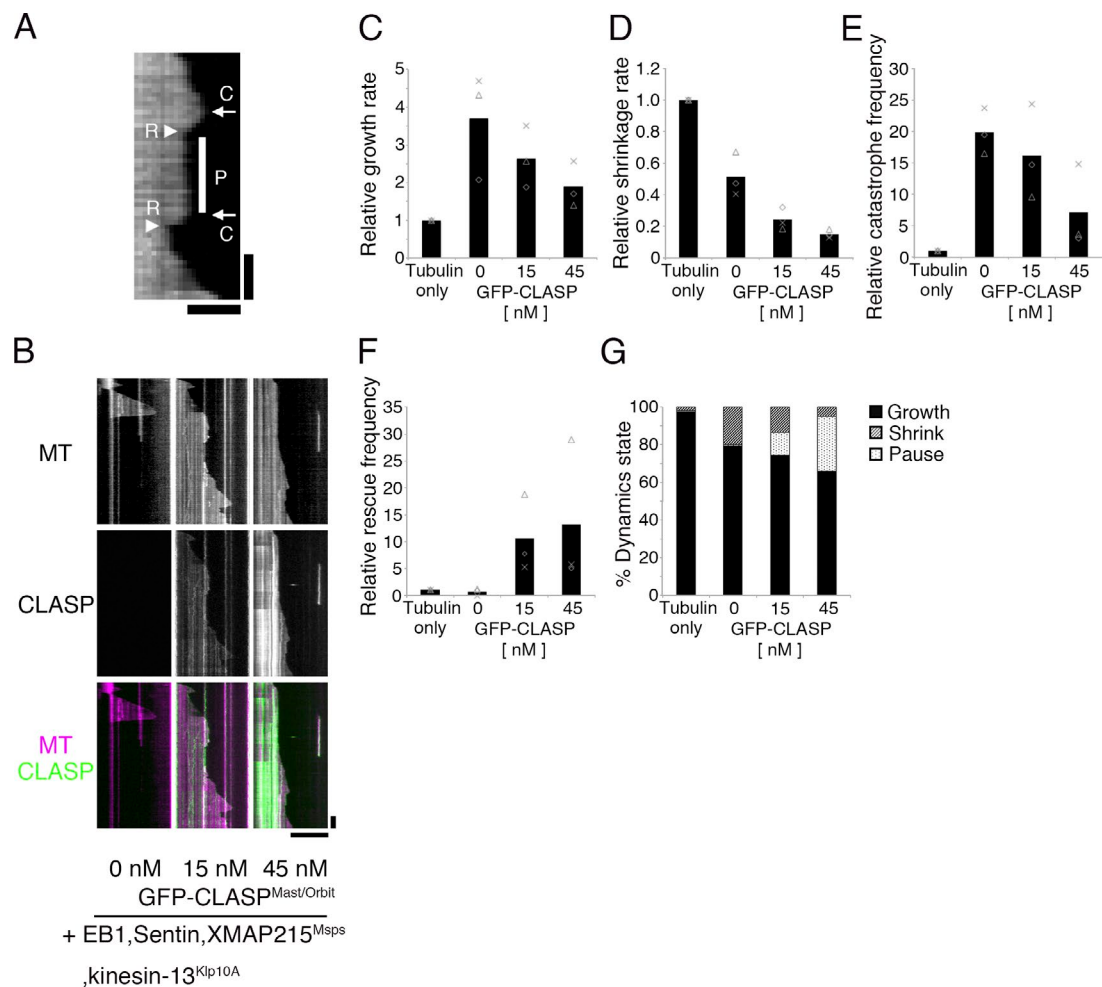


Figure 2. Reconstitution of dynamic MTs with five factors. (A) An example of pause (P) identified in the kymograph. Catastrophe (C) and rescue (R) events are also marked. Bars: (horizontal) 2 μ m; (vertical) 30 s. (B–G) Kymographs and kinetic parameters of MT polymerization dynamics with 0–45 nM GFP-CLASP^{Mast/Orbit} in the presence of 15 μ M S2 tubulin, XMAP215^{Mps}, EB1, Sentin, and kinesin-13^{Klp10A}. The mean values of each experiment are marked in gray, whereas the mean values of all the experiments are indicated by black columns (three independent experiments). Relative values are plotted in the graphs, whereas the actual values are presented in Table 1. Bars: (horizontal) 10 μ m; (vertical) 1 min.

Reconstitution of three-phase MT dynamics with five factors

We next mixed all five factors in the reaction with S2-derived tubulin. Interestingly, we observed that MTs repeated cycles of shrinkage and growth, accompanied by frequent rescue and pausing; this behavior had never been observed using just one

to three factors (Fig. 2 and Table 1). Each parameter value was within range of the corresponding cellular value, and the overall stochastic behavior was reminiscent of interphase MT dynamics in S2 cells (Fig. S2).

To validate the similarity between in vitro and in vivo dynamics, we omitted kinesin-13^{Klp10A}, Sentin, or CLASP^{Mast/Orbit}

Table 1. Kinetic parameters of in vitro-reconstituted MT polymerization dynamics with the addition of five factors

Condition	CLASP concentration	n	Growth rate	Shrinkage rate	Catastrophe frequency	Rescue frequency
	nM		μ m/min	μ m/min	$\times 10^{-3}$ s ⁻¹	$\times 10^{-3}$ s ⁻¹
Tubulin alone	0	3	1.6 \pm 0.1 (60)	37.9 \pm 8.2 (30)	0.9 \pm 0.2 (31)	10.5 \pm 6.1 (6)
Plus EB1/Sentin/MspS/Klp10A/CLASP	0	3	5.8 \pm 2.2 (88)	19.0 \pm 3.6 (86)	17.7 \pm 2.0 (86)	5.3 \pm 5.8 (7)
	15	3	4.2 \pm 1.3 (260)	9.1 \pm 2.5 (219)	13.9 \pm 3.6 (217)	81.6 \pm 11.2 (211)
	45	3	3.0 \pm 1.0 (166)	5.6 \pm 0.9 (88)	5.7 \pm 3.9 (86)	86.1 \pm 23.9 (87)
P (CLASP 45 nM vs. 0 nM)			0.116 (n = 3) 0.002 (exp1) <10 ⁻⁴ (exp2) <10 ⁻⁴ (exp3)	0.003 (n = 3) <10 ⁻⁴ (exp1) <10 ⁻⁴ (exp2) <10 ⁻⁴ (exp3)	0.009 (n = 3)	0.005 (n = 3)

Values are mean \pm SD of three independent experiments (total number of events). See Fig. 2 for a graphic presentation of these results.

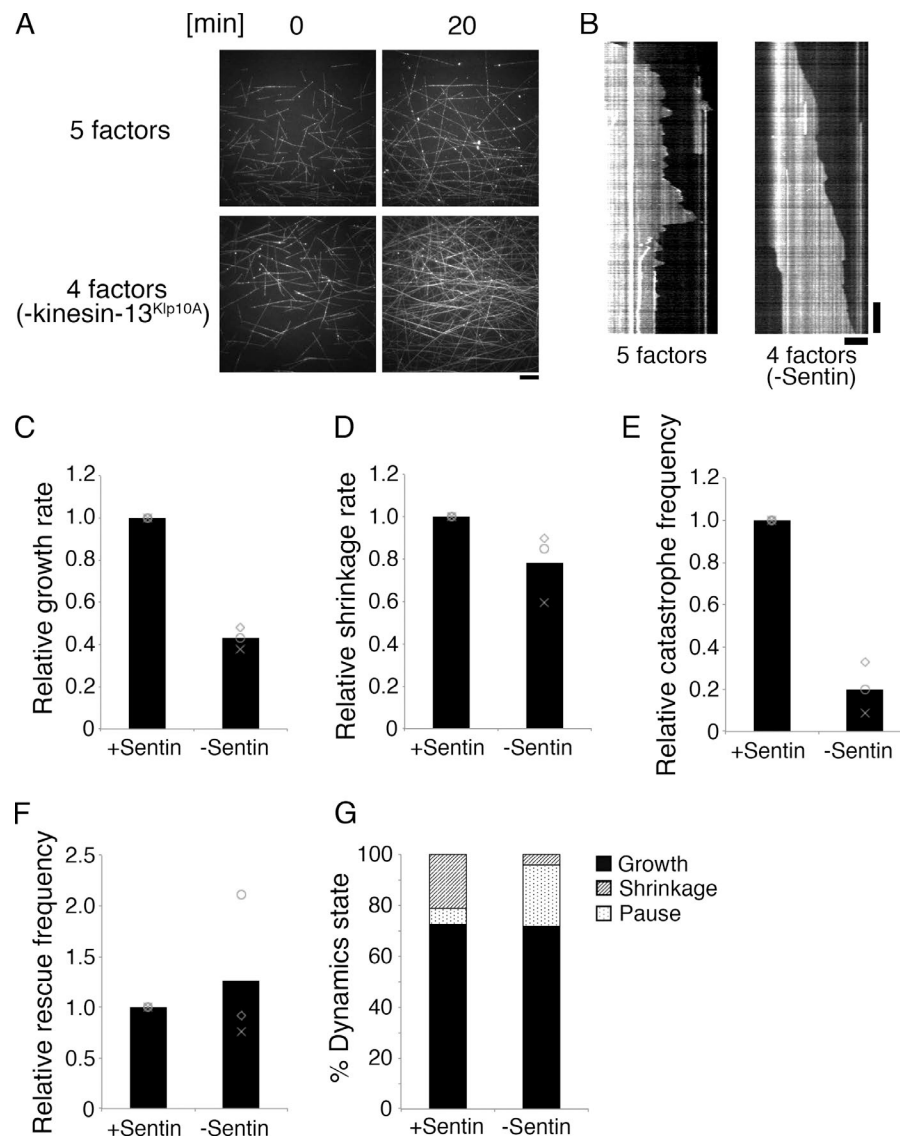


Figure 3. Effect of Sentin or kinesin-13^{Klp10A} removal from the MT polymerization reaction. (A) MTs dramatically increased because of reduced catastrophe frequency in the absence of kinesin-13^{Klp10A}. The other four factors were mixed with 15 μ M S2 tubulin for 20 min. Bar, 10 μ m. (B–G) MT polymerization dynamics in the presence or absence of Sentin. The other four factors (EB1, XMAP215^{Msps}, kinesin-13^{Klp10A}, and 15 nM CLASP^{Mast/Orbit}) were mixed with 15 μ M S2 tubulin (three independent experiments). Actual values are presented in Table S1. Bars: (horizontal) 5 μ m; (vertical) 2 min.

from the reaction and compared the outcome with *in vivo* RNAi results. In S2 cells, kinesin-13^{Klp10A} knockdown decreases the catastrophe frequency (Mennella et al., 2005). Sentin knockdown inhibits the dynamic nature of MTs, specifically through increasing the pause duration and decreasing growth and shrinkage rates and catastrophe frequency (Li et al., 2011). CLASP^{Mast/Orbit} depletion in S2 cells increased the growth and shrinkage rates but reduced the pause duration (Sousa et al., 2007; Trogden and Rogers, 2015). Interestingly, we generally reproduced these tendencies in our *in vitro* assays through removal of kinesin-13^{Klp10A}, Sentin, or CLASP^{Mast/Orbit}. Without kinesin-13^{Klp10A}, catastrophe was dramatically suppressed and the imaging field was occupied by numerous MTs (Fig. 3 A; only seven catastrophe events were observed during a 413-min growing phase). In the case of CLASP^{Mast/Orbit}, all seven parameters were changed in a qualitatively consistent manner (Fig. 2, B–F; “0 nM CLASP” corresponds to CLASP depletion; see also Fig. S3 [A–E], for comparison to published *in vivo* results). Sentin depletion had a similar impact; changes in six out of seven parameters were consistent *in vivo* and *in vitro* (Fig. 3, B–G; Fig. S3, F–J; and Table S1).

Lattice-bound CLASP^{Mast/Orbit} induces pausing in shrinking MTs

Because pausing was rarely observed in the absence of CLASP^{Mast/Orbit}, we investigated whether GFP-CLASP^{Mast/Orbit} accumulation is correlated with pause induction. Because CLASP^{Mast/Orbit} was often abundantly localized to the growing end of MTs via EB1 binding, we focused our analysis on shrinking MTs. We first surveyed GFP-CLASP^{Mast/Orbit} signal distribution along the lattice of shrinking MTs ($n = 21$) and observed that 24% of the pixels contained a high GFP signal (Fig. 4 A). This result indicated that if pausing is induced at random locations, the event has a 24% chance of occurring at GFP-accumulation sites. However, we identified 51% ($n = 51$) of shrinkage-to-pause events occurring at GFP-occupied sites (red in Fig. 4, B and C). Furthermore, in 9 out of the 25 events, in which pausing was induced at sites with low GFP-CLASP^{Mast/Orbit} abundance, MTs already had strong GFP signals during shrinking (green in Fig. 4, B and C). Thus, GFP-CLASP^{Mast/Orbit} accumulation and pause induction were well correlated, albeit not perfectly. To investigate whether CLASP^{Mast/Orbit} on the lattice recruits other factors, we mixed MT seeds with

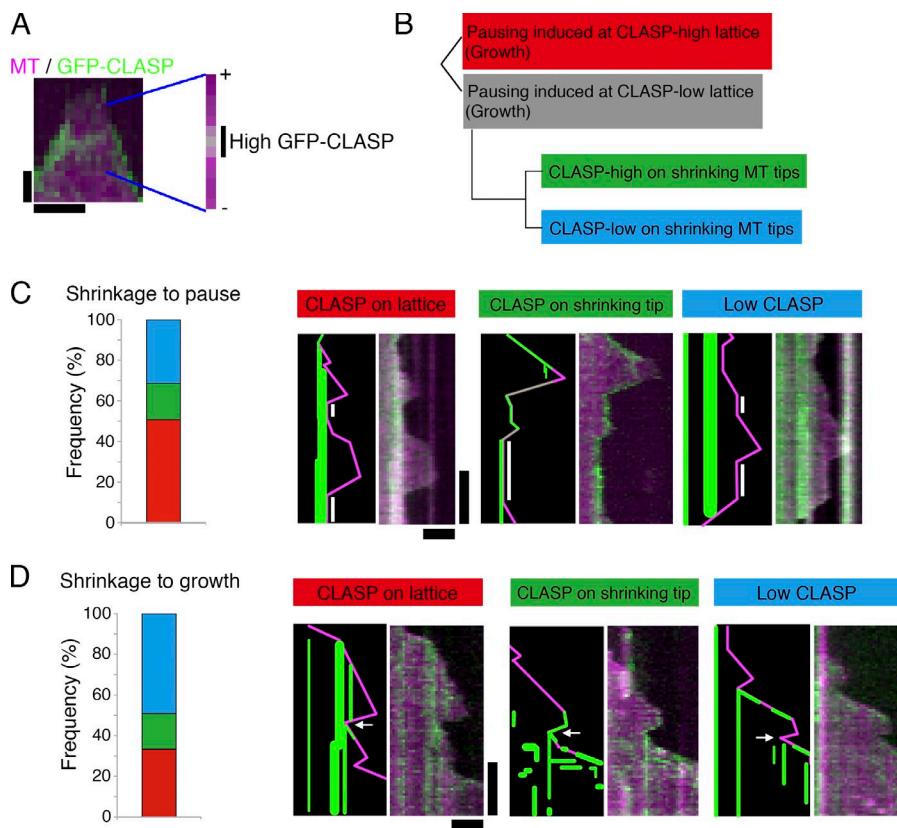


Figure 4. Pausing was induced at the CLA SP^{Mast/Orbit} accumulation site with a high probability. (A) GFP-CLASP^{Mast/Orbit} distribution along the MT. An MT that just started to shrink is highlighted. Marked are the regions at which GFP-CLASP^{Mast/Orbit} was accumulated. Bars: (horizontal) 30 s; (vertical) 1 μ m. (B–D) Classification of shrinkage-to-pause (C; $n = 51$) or shrinkage-to-growth (D; $n = 63$) events based on GFP-CLASP^{Mast/Orbit} accumulation at the turning point. Kymographs and the schematic drawings are displayed. Actual data are derived from the experiments performed in Fig. 2 (shrinkage to growth) or Fig. 5 (shrinkage to pause). Pausing phases are marked with white bars, whereas arrows indicate the rescue points for regrowth. Bars: (horizontal) 2 μ m; (vertical) 1 min.

TagRFP-CLASP^{Mast/Orbit} and GFP-Sentin; Sentin is proposed to act as a bridge between CLASP^{Mast/Orbit} and XMAP215^{Msp} (Trogdon and Rogers, 2015) but does not localize to the MT lattice on its own (Li et al., 2012). We did not observe lattice localization of the GFP-Sentin signals in the presence of TagRFP-CLASP^{Mast/Orbit}, suggesting that prerecruitment of the Sentin–XMAP215^{Msp} complex is not required for CLASP^{Mast/Orbit} to induce pausing (Fig. S1 H).

Similar analysis was performed for shrinkage-to-growth events (Fig. 4, B and D). In this case, the correlation was weaker than for the pause but was still higher than expected from random occurrence (33% events occurred at GFP accumulation site [$n = 63$], which occupied 15% of the pixels [$n = 31$]). This result indicates that small amounts of CLASP^{Mast/Orbit} can also trigger the shrinkage-to-growth rescue, which differs from the results of fission yeast CLASP^{cls1}, wherein enrichment was perfectly correlated with the rescue event (Al-Bassam et al., 2010). We could not rule out the possibility that our microscope was not sensitive enough to detect moderately accumulated GFP-CLASP^{Mast/Orbit}. Regardless, the results suggest that fly CLASP is a potent pause/rescue-inducing factor in the presence of four other factors.

Increase in MT dynamicity by Plk1^{Polo} kinase

MT dynamics are subjected to mitotic regulation. During mitosis, the rate of astral MT growth and the frequency of catastrophe events increase, whereas the frequency of rescue events decreases (Belmont et al., 1990; Rusan et al., 2001). It was shown that the addition of active Cdk1 to the interphase frog extracts triggers the interphase-mitosis transition of MT dynamics (Verde et al., 1990). However, the event downstream of Cdk1 remains unclear. We first hypothesized that Cdk1 might

directly phosphorylate one or more of the five factors, or tubulin itself, to modulate MT dynamics. Autoradiography experiments indicated that kinesin-13^{Klp10A} and XMAP215^{Msp} are phosphorylated by the recombinant Cdk1–cyclin B complex (Fig. S4 A). Next, we incubated the five proteins and tubulin with Cdk1–cyclin B, followed by a MT polymerization assay. Here, kinetic parameters were not significantly or reproducibly changed in four experiments comparing the effect of Cdk1 (Fig. S4, B–F; and Table S2). These results were not consistent with the idea that Cdk1 phosphorylation directly regulates the activities of these six proteins, although they did not entirely disprove it.

We considered it likely that a factor other than Cdk1 is the direct contributor to the increase in MT dynamicity during mitosis. Polo-like kinase 1 (Plk1) consists of a complex regulatory circuit during mitotic entry (Lindqvist et al., 2009; Zitouni et al., 2014). We therefore next hypothesized that Plk1 may be responsible for the change in dynamics. Interestingly, incubation of five factors and tubulin with recombinant Plk1^{Polo} affected kinetic parameters in a reproducible manner; growth rate and catastrophe increased, whereas rescue and pausing were reduced, making MTs more dynamic (Fig. 5, B and C; and Table S3). Phos-tag and autoradiography analyses showed that kinesin-13^{Klp10A}, Sentin, and CLASP^{Mast/Orbit} were phosphorylated by Plk1^{Polo}, whereas no gel shift was observed after Phos-tag analysis for EB1, XMAP215^{Msp}, or S2 tubulin (Fig. 5 A). When GFP-CLASP^{Mast/Orbit} was mixed with tubulin, we observed a reduction in MT lattice association in the presence of Plk1^{Polo} (Fig. 5, D and E). This result was consistent with the increase of MT dynamicity conferred by Plk1^{Polo}, as lattice-associated CLASP^{Mast/Orbit} frequently induced pausing (Fig. 4 C). This effect was at least in part phosphorylation dependent, because kinase-inactive (kinase-dead [KD]) Plk1^{Polo}, purified in an identical manner to wild type, showed a milder effect (Fig. 5, D

and E). Conversely, when active $\text{Plk1}^{\text{Polo}}$ was mixed with the other four factors and tubulin, we observed rapid depolymerization of MT seeds (Fig. 5, F and G). The seed depolymerization suggested that kinesin-13 $^{\text{Klp10A}}$ is activated by $\text{Plk1}^{\text{Polo}}$. Thus, $\text{Plk1}^{\text{Polo}}$ -mediated phosphorylation affected the activities of CLASP^{Mast/Orbit} and kinesin-13 $^{\text{Klp10A}}$. Whether Sentin, which also has catastrophe-inducing activity (Li et al., 2012), is a critical substrate of $\text{Plk1}^{\text{Polo}}$ remains to be determined.

To investigate the involvement of $\text{Plk1}^{\text{Polo}}$ in MT dynamics in vivo, we measured growth velocity of mitotic astral MTs in S2 cells in the presence or absence of the $\text{Plk1}^{\text{Polo}}$ inhibitor BI2536. S2 cells have heterogeneity in centrosome numbers, which might alter the parameters associated with MT dynamics of each aster. Therefore, we measured astral MT dynamics in the monopolar spindle that was induced by RNAi knockdown of the kinesin-5 motor Klp61F (Fig. 6 A). Interestingly, mitotic EB1-GFP velocity was decreased by 40% after BI2536 treatment, which matched the velocity in interphase (Fig. 6 B). The decrease was unlikely induced by off-target effect of BI2536 on tubulin, because the MT growth rate during interphase was unaffected by this treatment. We next used the same dataset to compare growth time. Because pausing is a rare event during S2 mitosis (Li et al., 2011), we considered that disappearance of the EB1-GFP signals in the mitotic aster mostly corresponded to catastrophe. Duration of EB1-GFP tracking was 56% longer in the presence of BI2536, and this was similar to interphase values, suggesting that $\text{Plk1}^{\text{Polo}}$ elevated catastrophe frequency in mitotic cells (Fig. 6 C).

Discussion

This study aimed to reproduce cellular MT polymerization dynamics in vitro more faithfully than previous studies. Toward this goal, we expanded the materials for reconstitution to seven purified proteins (including tubulin and a kinase). To the best of our knowledge, this presents the most complex reconstitution of MT polymerization dynamics. Our in vitro experiments, together with previous loss-of-function analyses in vivo, suggest that five proteins, XMAP215 $^{\text{Msps}}$, EB1, Sentin, kinesin-13 $^{\text{Klp10A}}$, and CLASP^{Mast/Orbit}, constitute the core machinery that regulates MT dynamics in the S2 cytoplasm (Fig. 6 D, left). In the presence of these five factors, typical MT behavior was reproduced, in which growth, shrinkage, and pausing phases repeat in a stochastic order. The change in dynamics in the absence of CLASP^{Mast/Orbit}, Sentin, or kinesin-13 $^{\text{Klp10A}}$ supports this model. Omission of each factor in vitro resulted in MT dynamics that deviated from that of untreated S2 cytoplasm; dynamics became more similar to that observed upon RNAi depletion of corresponding factors. Moreover, phosphorylation by the mitotic kinase $\text{Plk1}^{\text{Polo}}$ resulted in more dynamic MTs, reminiscent of what occurs during interphase-to-mitosis transitions (Fig. 6 D, right).

CLASP induces pausing

A previous study using purified yeast CLASP^{Clsp1} identified rescue (shrinkage-to-growth) and catastrophe-inhibiting activities, but not pausing or growth-inhibiting activity (Al-Bassam et al., 2010). An intriguing observation in the current study was the emergence of pause when CLASP^{Mast/Orbit} was added with the four other factors. This illustrates the value of multifactor reconstitution and suggests that an equilibrium of CLASP^{Mast/Orbit}

with MT polymerase and depolymerase produces such a state. It is possible that such activity was conferred by physical interaction with other factors; for instance, it was recently shown that XMAP215 $^{\text{Msps}}$ interacts with CLASP^{Mast/Orbit} via Sentin in S2 cells (Trogden and Rogers, 2015). In S2 cells, RNAi depletion of EB1, Sentin, or XMAP215 $^{\text{Msps}}$ increases the pause duration in interphase, and this phenotype could not be readily explained (Rogers et al., 2002; Brittle and Ohkura, 2005; Li et al., 2011). On the basis of current in vitro results, this phenotype may be interpreted that when EB1, Sentin, or XMAP215 is depleted, CLASP^{Mast/Orbit} dominates at the plus end and induces pausing together with the other three factors.

Plk1 elevates MT dynamicity

Another finding in this study was the increase in MT dynamicity by $\text{Plk1}^{\text{Polo}}$ kinase. The involvement of $\text{Plk1}^{\text{Polo}}$ is not inconsistent with the classical analysis in which Cdk1 triggers elevation of MT dynamicity in frog egg extracts (Verde et al., 1990). It has been shown that Cdk1 and Plk1 comprise a complex signaling network, and the addition of Cdk1 to egg extracts may well activate the Plk1 kinase. Consistent with the in vitro data, we observed reversion of MT dynamics, from mitotic to interphase mode, by treating prometaphase cells with a $\text{Plk1}^{\text{Polo}}$ inhibitor. An in vitro kinase assay showed that Sentin, kinesin-13 $^{\text{Klp10A}}$, and CLASP^{Mast/Orbit} are the substrates of $\text{Plk1}^{\text{Polo}}$. The appearance of multiple bands in the electrophoresis gel indicates that multiple residues are potential $\text{Plk1}^{\text{Polo}}$ phosphorylation sites. Determining the substrate residues that are phosphorylated to influence MT dynamics in vivo is an important future research topic. However, our results do not exclude the possibility that $\text{Plk1}^{\text{Polo}}$ phosphorylation of other proteins also contributes to changes in MT dynamics in cells.

Comparison between in vitro and in vivo data

Although we believe we have achieved hitherto the most in vivo-relevant reconstitution, we admit that the study is still not a precise reproduction of S2 MT dynamics. First, other factors may also contribute to MT dynamics in cells. S2 cells express additional factors that can modulate MT dynamicity in vitro, such as D-TPX2 and two other kinesin-13 family members (Klp59C and Klp59D); however, RNAi depletion of these factors results in a phenotype that is not as severe as that observed after ablation of the five factors used in this study (Goshima and Vale, 2003; Mennella et al., 2005; Goshima, 2011). Second, our data on MT dynamics are basically limited to free MTs in the cytoplasm. Other types of MTs, such as cortical and peripheral MTs, are often bound to other protein or membranous structures and show different dynamic kinetics (Trogden and Rogers, 2015), and additional factors are likely necessary for reconstitution. For instance, human CLASP depletion decreased catastrophe as well as rescue at the cell cortex, which is not entirely consistent with the consequence we observed here in vitro (Mimori-Kiyosue et al., 2005); GSK3 kinase is known to phosphorylate CLASP and thereby regulate peripheral MT dynamics (Kumar et al., 2009; Trogden and Rogers, 2015). Kinetochore MTs in metaphase are also bound to many other proteins. However, a key feature of kinetochore MT dynamics might have been partially reproduced with one of our assay conditions. In cells, CLASP^{Mast/Orbit} is most prevalent at the outer kinetochore region during metaphase and is required for kinetochore MT polymerization (Maiato et al., 2005). The slow but persistent growth of

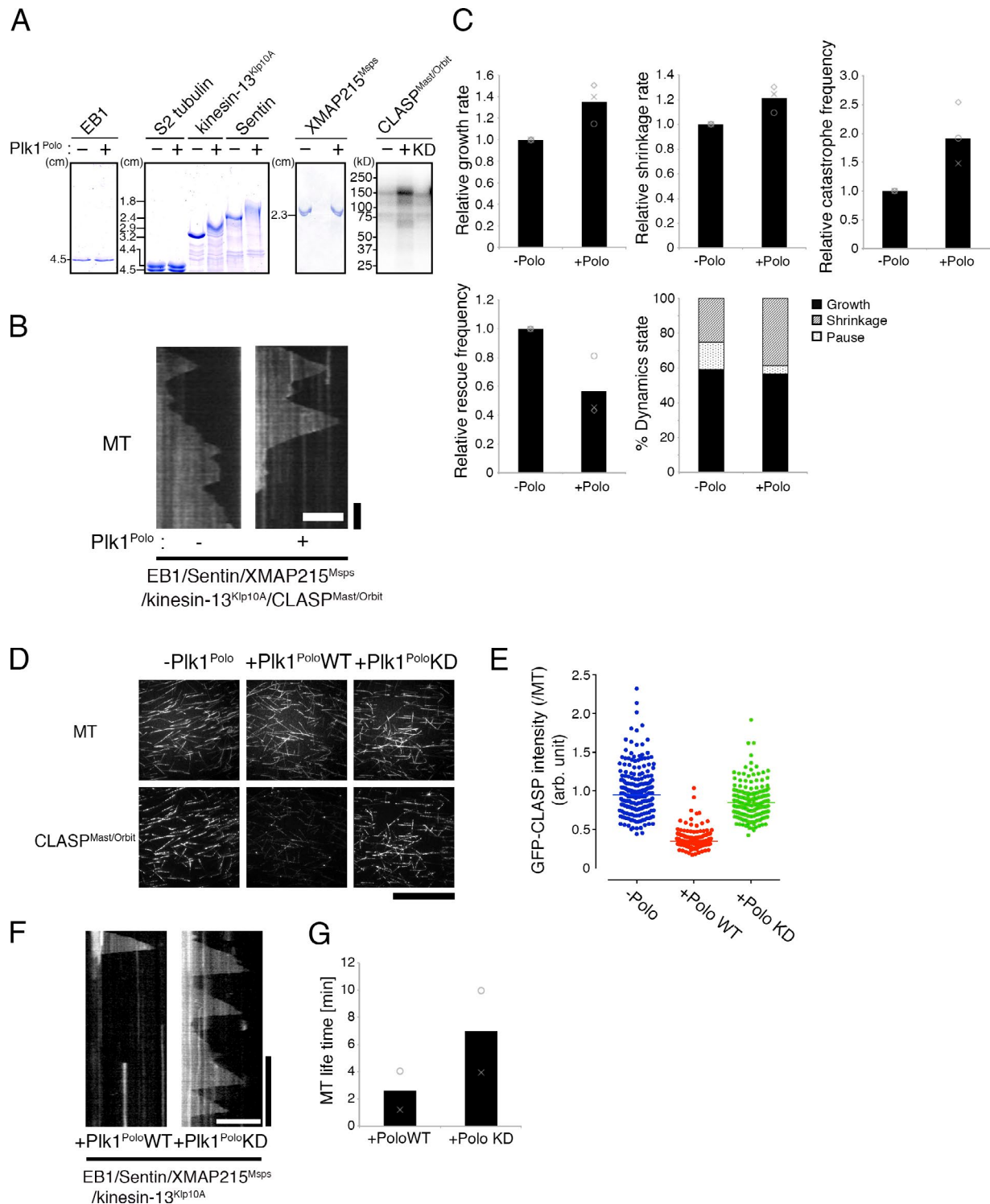


Figure 5. Phosphorylation by $\text{Plk1}^{\text{Polo}}$ elevated MT dynamicity in vitro. (A) Autoradiography (rightmost panel) and Phos-tag gel electrophoresis (others) showed phosphorylation of $\text{CLASP}^{\text{Mast/Orbit}}$, kinesin-13 Kip10A , and Sentin by $\text{Plk1}^{\text{Polo}}$. The markers of the Phos-tag gel indicate the distance from the top of the resolving gel. (B and C) Kymographs and kinetic parameters showing that treatment of five MT dynamic-modulating factors with $\text{Plk1}^{\text{Polo}}$ makes MTs more dynamic (10 μM S2 tubulin; 45 nM $\text{CLASP}^{\text{Mast/Orbit}}$). In the graph, the mean and relative values of each experiment are plotted (three independent experiments). Actual values are presented in Table S3. Bars: (horizontal) 5 μm ; (vertical) 1 min. (D and E) Association of 5 nM GFP- $\text{CLASP}^{\text{Mast/Orbit}}$ with the MT lattice (10 μM S2 tubulin) was drastically reduced by incubation with 100 nM wild-type $\text{Plk1}^{\text{Polo}}$ ($P < 0.0001$, Mann-Whitney U test) and more modestly with a kinase-dead (KD) mutant ($n = 227$ [-Polo], 212 [+PoloWT], and 214 [+PoloKD]). Bar, 50 μm . (F and G) MT depolymerization activity of kinesin-13 Kip10A was enhanced by incubation with wild-type $\text{Plk1}^{\text{Polo}}$ (two independent experiments). Four factors (EB1, XMAP215 $^{\text{Msps}}$, Sentin, and 10 nM kinesin-13 Kip10A) and 15 μM S2 tubulin were included in this reaction, and seed depolymerization was assessed after generating the kymograph. Note that the lifetime for wild-type $\text{Plk1}^{\text{Polo}}$ is likely overestimated because some seeds had been already depolymerized when the imaging was started. Bars: (horizontal) 10 μm ; (vertical) 5 min.

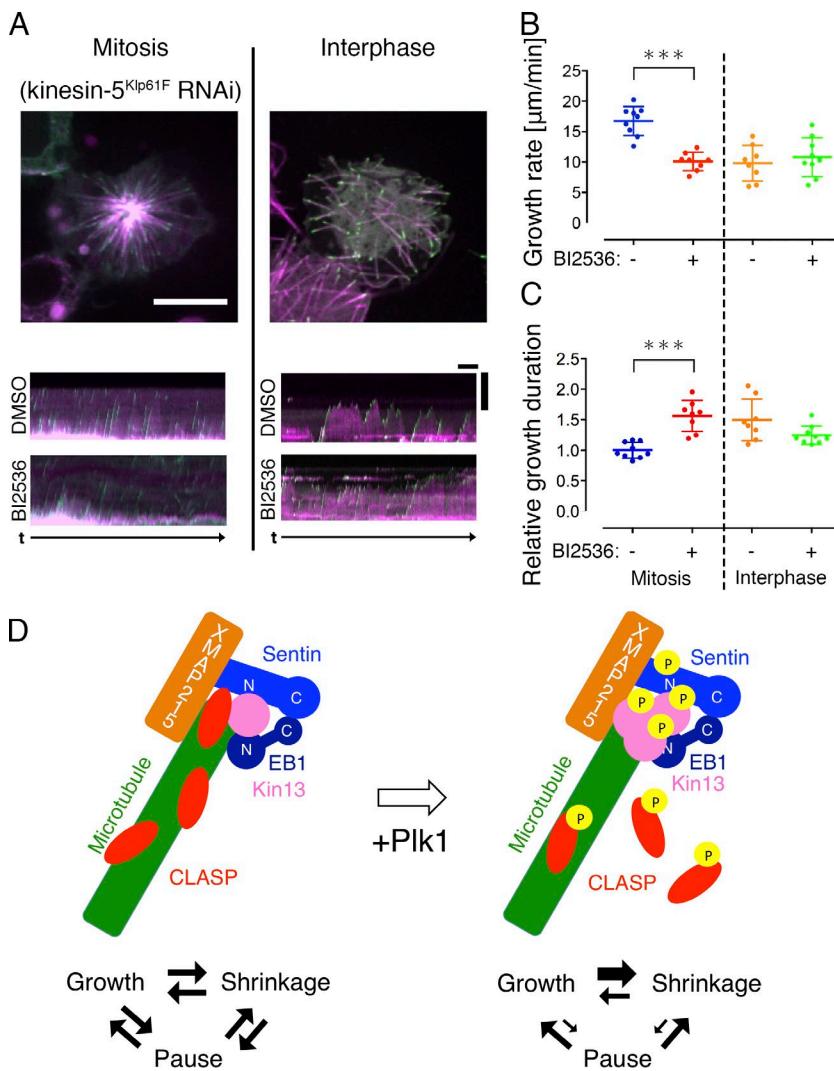


Figure 6. $\text{Plk1}^{\text{Polo}}$ inhibition dampens MT dynamics in vivo. (A–C) $\text{Plk1}^{\text{Polo}}$ inhibition decreased MT growth rate and catastrophe frequency in mitosis, but not in interphase. Mitotic astral MTs in the monopolar spindle induced by Klp61F (kinesin-5) RNAi were analyzed (100–200 comets per cell, and eight or nine cells in each condition). Each dot in the graph represents the mean value obtained from a cell. Asterisks indicate statistically significant differences (***, $P < 0.0001$; Turkey's multiple comparisons test; the normality of data distribution was checked by D'Agostino-Pearson omnibus normality test and Shapiro-Wilk normality test). Error bars indicate SDs. Bar for cell images, 10 μm ; horizontal bar for kymographs, 1 min; vertical bar for kymographs, 5 μm . (D) Reconstitution of regulated microtubule polymerization dynamics was achieved in this study. Three-phase polymerization cycles were observed when five factors were mixed with tubulin. Addition of the active Plk1 kinase increased MT dynamicity, wherein pausing became rare.

kinetochore MTs has resemblance to that observed when a high concentration of CLASP^{Mast/Orbit} was mixed with four other factors; MTs grew slower and rarely shrank even in the presence of catastrophe-promoting factors (Fig. 2; 45 nM CLASP^{Mast/Orbit}). Thus, although multiple other MT-regulating proteins are present at the outer kinetochore, it is tempting to speculate that accumulation of CLASP^{Mast/Orbit} is a key event that distinguishes kinetochore MTs from highly dynamic astral MTs.

Finally, another difference between our system and the *in vivo* situation involves the environment and protein concentrations. We used a standard nonviscous buffer in our assay, which would be different from the cytoplasm that is highly viscous, containing many other molecules. Furthermore, our rough estimate indicated that cellular concentrations of CLASP^{Mast/Orbit} and kinesin-13^{Klp10A} were 45- to 80-fold higher than what we used *in vitro*, a level currently unachievable by our purification method (Fig. S5). One possible explanation for this discrepancy is that cellular proteins are regulated spatially or posttranslationally, and the number of actual proteins that can target MTs is small. Alternatively, the antagonistic activity of CLASP^{Mast/Orbit} and kinesin-13^{Klp10A} on catastrophe suggests that the molecular ratio of the two proteins rather than the absolute abundance of each protein might be critical. In support of this idea, the ratio of CLASP^{Mast/Orbit} and kinesin-13^{Klp10A} was comparable between

in vitro (1:6.7) and *in vivo* (1:3.7) conditions. Regardless, our reconstitution admittedly is qualitative, and quantitative comparison with *in vivo* kinetic parameters remains difficult. Converting the kinetic parameters and protein concentrations to *in vivo* values remains a key challenge in the reconstitution of MT dynamics and perhaps also for other complex cellular events (Shintomi et al., 2015).

Toward reconstitution of mammalian MT dynamics

The data obtained in this study might be largely applicable to mammalian cells. EB1, XMAP215^{MspS}, kinesin-13^{Klp10A}, CLASP^{Mast/Orbit}, and $\text{Plk1}^{\text{Polo}}$ have clear orthologues in mammals. Although not conserved in amino acid sequences, Sentin is likely a functional homologue of mammalian SLAIN2, because both proteins bind to EB1 and XMAP215^{MspS}, bridging these two molecules for synergistic MT growth acceleration (Li et al., 2011; van der Vaart et al., 2011). Furthermore, Plk1 -dependent activation of kinesin-13 has been demonstrated *in vitro* (Jang et al., 2009; Zhang et al., 2011). Because multiple paralogous/homologous genes have been identified for CLASP (1 and 2), EB (1, 2, 3), SLAIN (1 and 2), and kinesin-13 (Kif2a, Kif2b and MCAK), mammalian cells might have further layers of complexity compared with *Drosophila*. In fact, it was recently

shown that human CLASP1 and an isoform of CLASP2 have common and also distinct activities (Yu et al., 2016). Regardless, a reasonable first attempt toward reconstitution of mammalian MT dynamics would be to purify and mix homologues of the five proteins identified in this study.

Materials and methods

Cloning

Primers to amplify full-length CLASP^{Mast/Orbit} were 5'-CACCAGGG CTATCGGAAGCCCAGCGACCTGG-3' and 5'-TGACGACGATGC CGCGGAGGAGTTCTTGG-3'. The amplified DNAs were cloned into pENTR-D/TOPO, digested with NotI and AscI, blunted, and cloned into the pFastBac-His-mGFP vector (StuI digested). His-CLASP^{Mast/Orbit} was constructed by Gateway cloning into the pDEST10 vector. Kinesin-13^{Klp10A} gene was amplified with the primers 5'-CACCAGGG CATGATTACGGTGGGGCA-3' and 5'-CTAACGCTTGCCATTCCGG CGAA-3'. The amplified DNAs were cloned into pENTR-D/TOPO, followed by Gateway cloning into pDEST17. Plk1^{Polo} was amplified with 5'-TTGCGGCCGCCACCATGGCCGCGAAGCCCCGAGG-3' and 5'-TTGGCGCGCCTATGTGAACATCTCTCCAGC-3', and ligated into pENTR-D/TOPO after NotI-AscI digestion. The gene was then cloned into pDEST10 by the Gateway reaction.

Protein purification

S2 tubulin was purified by using a previously described method (Widlund et al., 2012). GST-tagged TOG1 domain (*Saccharomyces cerevisiae* Stu2, 1–306 aa) expression was induced in *E. coli* BL21 in 2 liters of Terrific Broth, 0.2 mM IPTG for 18 h at 25°C (TOG1-encoding plasmid was a gift from P. Widlund, Max Planck Institute, Dresden, Germany). Harvested cells were lysed with 120 ml 2×PBS (5.4 mM KCl, 3 mM KH₂PO₄, 16.2 mM Na₂HPO₄, and 274 mM NaCl) supplemented with 500 U benzonase (EMD Millipore) and protease inhibitors (0.5 mM PMSF and peptide cocktails [1 µg/ml leupeptin, pepstatin, chymostatin, and aprotinin]) by the Advanced Digital Sonifier D450 (Branson). The lysate was incubated with 5 ml GSTrap (GE Healthcare) for 1 h at 4°C. After washing once, GST-TOG1 were eluted with 20 mM reduced glutathione in 2×PBS, pH 8.0, followed by dialysis using PD-10 desalting column (GE Healthcare) and 3.5 ml coupling buffer (100 mM NaHCO₃ with 100 mM NaCl at pH 8.2). After dialysis, 80 mM MgCl₂ was added. For conjugation of GST-TOG1, 3.5 ml GST-TOG1 protein solution was applied to 1 ml HiTrap NHS-activated HP columns (GE Healthcare; preactivated by 1 mM HCl) and recirculated at 1.5 ml/min in the column for 20 min. The remaining reactive groups were blocked by a 30-min incubation with 0.5 M ethanolamine and 0.5 M NaCl, pH 8.3. The column was then washed with 1×PBS containing 50% glycerol and was then stored at –20°C. 700 ml S2 cell culture was lysed with 7 ml MRB80 buffer (80 mM K-Pipes, pH 6.8, 4 mM MgCl₂, and 1 mM EGTA) supplemented with 125 U benzonase (EMD Millipore) by a Dounce tissue grinder. The lysate was clarified by 9,400 g and 135,000 g centrifugation and applied to a 1-ml TOG1-conjugated column. Tubulin was eluted with the elution buffer (0.5 M (NH₄)₂SO₄ in MRB80, 10 µM GTP), followed by desalting with the PD-10 desalting column (GE Healthcare) and MRB80. The protein solution was concentrated to >40 µM by Amicon Ultra-4 MWCO 30K. The concentrated solution was flash frozen with liquid nitrogen and stored at –80°C. EB1 was purified using *E. coli* following a protocol described previously (Li et al., 2011, 2012). GST-EB1 expression was induced in BL21-AI with 0.2% arabinose for 16 h at 25°C. Harvested cells were lysed using the Advanced Digital Sonifier D450 (Branson) and the lysis buffer (25 mM Tris-HCl, 1 mM EGTA,

150 mM NaCl, and 1 mM DTT) supplemented with protease inhibitors (0.5 mM PMSF and peptide cocktails). After incubation with glutathione Sepharose beads for 1 h at 4°C, GST was cleaved using HRV 3C protease (Thermo Fisher Scientific), followed by gel filtration chromatography (MRB80, 100 mM KCl, and 1 mM DTT). His-kinesin-13^{Klp10A} was purified using the standard Ni-NTA purification with the buffer identical to that of EB1, except that 1 mM ATP was supplied. His-Sentin, His-GFP-Sentin, and XMAP215^{Msp}-HAHis were purified with insect SF21 cells essentially as described previously (Li et al., 2012). 1.5 × 10⁶ uninfected SF21 cells ml^{–1} in an Erlenmeyer flask were treated with baculovirus and incubated with shaking for 3 d (120 rpm at 27°C). The cells were lysed using 5 ml (per 50-ml cell culture) lysis buffer (50 mM K-Hepes, 100 mM KCl, 1 mM MgCl₂, 1 mM EGTA, 1 mM DTT, 30 mM imidazole, and 1% Triton X-100, pH 7.6, with protease inhibitors). After incubation with nickel-coated beads (0.5-ml bed volume) for 1 h, the proteins were eluted using 250 µl elution buffer (lysis buffer supplemented with 200 mM imidazole without 1% Triton X-100). Elution was repeated five to seven times. Buffer was exchanged to 1× MRB80 containing 100 mM KCl and 1 mM DTT using the desalting column PD MiniTrap G-25 (GE Healthcare). His-GFP-CLASP^{Mast/Orbit} (wild type and TOG mutant [W334E, K421A, W852E, K940A, which reside in the TOG2 and TOG3 domains]), His-CLASP^{Mast/Orbit}, TagRFP-CLASP^{Mast/Orbit}, and His-Plk1^{Polo} (wild type and K54A KD mutant) were purified in a manner identical to that used for Sentin, except that the lysis buffer contained 100 mM KCl and 0.1–0.2% Tween-20 (CLASP), or cells were treated with 100 nM okadaic acid 1 h before collection (Plk1^{Polo}). Purified S2 tubulin, EB1, kinesin-13^{Klp10A}, and Plk1^{Polo} were flash frozen (with 20% glycerol for EB1 and 20% sucrose for kinesin-13^{Klp10A} and Plk1^{Polo}), whereas other proteins were kept on ice and used within 48 h. Recombinant Cdk1–cyclin B complex was purchased from New England Biolabs, Inc. Gel filtration chromatography of GFP-CLASP^{Mast/Orbit} was performed with a Superdex 200 increase 10/300 GL column equilibrated with a gel filtration buffer (80 mM K-Pipes, pH 6.8, 100 mM KCl, 4 mM MgCl₂, 1 mM EGTA, and 1 mM DTT).

In vitro MT polymerization assay

The in vitro MT polymerization assay was performed basically following the method described in Li et al. (2012), which referred to Bieling et al. (2010) and Gell et al. (2010). Flow chambers (22 mm [width] × 1 mm [height] × 0.15 mm [depth]) were assembled between a coverslip and a precleaned micro slide glass with double-sided tape. Glass was soaked for 3 d in water supplied with the detergent SCAT 20-X, rinsed with fresh water, fired with methanol, and then silanized. The silanized coverslip was coated with antibiotic (1–5% in 1× MRB80; Invitrogen), and the nonspecific surface was blocked with Pluronic F127 (1% in 1× MRB80; Invitrogen). Biotinylated MT seeds (50–100 µM tubulin mix containing 10% biotinylated pig tubulin and 10% Alexa Fluor 568–labeled pig tubulin with 1 mM GMPPCP) were specifically attached to the functionalized surface by biotinylated tubulin–antibiotic links. After the chamber was washed with 1× MRB80, MT growth was initiated by flowing tubulin (containing 3.3% Alexa Fluor 568–labeled pig tubulin), EB1 (400 nM), Sentin (200 nM), GFP-CLASP^{Mast/Orbit} (0–45 nM), kinesin-13^{Klp10A} (100 nM, with 1 mM ATP), and/or XMAP215^{Msp}-HA (100 nM) into the assay buffer (1× MRB80, 75 mM KCl, 1 mM GTP, 0.5 mg/ml k-casein, and 0.1% methylcellulose) and an oxygen scavenger system (50 mM glucose, 400 µg/ml glucose-oxidase, 200 µg/ml catalase, and 4 mM DTT). The samples were sealed with candle wax. We used fixed concentrations of EB1, Sentin, and XMAP215^{Msp} based on our previous study, where the effect of these three factors combined has been demonstrated at these concentrations (Li et al., 2012). Kinesin-13^{Klp10A} was used at 100 nM, because increasing this protein

further depolymerized virtually all MT seeds in the imaging field. To match the estimated cellular tubulin concentration (Fig. S5), we used 15 μ M S2 tubulin for our main reconstitution experiments involving five or four nontubulin proteins (Figs. 2 and 3). However, in some experiments, we used 10 μ M S2 tubulin to save the protein whose yield was low compared with yields of pig brain-derived tubulin (Fig. 1 B and Fig. 5, B and D). When the samples were treated with Cdk1–cyclin B or Plk1^{Polo} kinase, 5 or 100 nM recombinant kinase (with 2 mM ATP), respectively, was added to the reaction. During the experiments, the samples were kept at 27°C; images were collected every 3 s for 20 min via total internal reflection fluorescence microscopy with a Ti system (Nikon), EMCCD camera (Evolve; Roper), 100 \times (1.49 NA) lens, and a 488-nm/561-nm excitation laser. The microscopes were controlled by Micromanager.

Data analysis

All parameters of MT plus end dynamics in vitro, except those related to pause, were determined in accordance with a method described previously (Li et al., 2012). In brief, kymographs were generated for every MT seed identified in the image field, and \sim 10 MTs, with a traceable plus end, were randomly selected. The starting and end points of growth and shrinkage were identified in the kymograph. The duration of and change in MT length for each phase were measured, and the rate was determined by calculating the mean velocity between the start and end points of each phase. Catastrophe frequency was determined by dividing the number of shrinkage events by the sum of growth and pause time. The transition from shrinkage to pause or growth was considered a rescue event, and the rescue frequency (for shrinkage time) was calculated. When MTs did not grow or shrink more than two pixels (\sim 0.35 μ m) for five or more frames (15 s), this period was defined as pause; this definition was comparable to that used in the in vivo pause measurement of our previous study (Li et al., 2011). In the analysis presented in Figs. 1 and S1, we did not count the pause phase, because it was impossible to identify the starting point of the pause phase of these slowly and persistently growing MTs; for them, we obtained the growth rate throughout. Catastrophe frequency in the absence of kinesin-13^{Plk10A} was measured by directly tracing the MT plus ends, because MTs were overly crowded and kymographs could not be generated.

Data presentation

We followed Li et al. (2012) for data presentation. As was described previously (Li et al., 2012), some degree of variation in MT behavior was observed on different experiment days, possibly because of differences in protein batches. To adjust for this variation, we have provided most of the graphs for the values for each condition relative to the control data acquired on the same day. In contrast, the mean values of three experiments are provided in the tables. For example, when we obtained growth rates of a_1 , a_2 , and a_3 (micrometers per minute) in three independent experiments using tubulin alone, as well as b_1 , b_2 , and b_3 (micrometers per minute) after CLASP^{Mast/Orbit} addition, we present the relative values as b_1/a_1 , b_2/a_2 , and b_3/a_3 in gray and $(b_1/a_1 + b_2/a_2 + b_3/a_3)/3$ in black in the figures. In the tables, however, we provide the mean rates of $(a_1 + a_2 + a_3)/3$ and $(b_1 + b_2 + b_3)/3$ separately with SDs. For all parameters, the p-values based on an unpaired two-sided Student's *t* test analysis of three or four experiments were obtained and are presented in Tables 1, S1, S2, and S3. However, the statistical significance of these differences as well as normality of the data distribution could not be properly discussed because sample sizes were too small; nevertheless, the values served as a good indicator of the degree of change. For growth and shrinkage rates where sufficiently large numbers of MTs were analyzed, we additionally provided p-values for each experiment

based on a Mann-Whitney *U* test (therefore, three or four p-values are displayed in each column).

Cell biology

S2 cells that express EB1-GFP and mCherry-tubulin were cultured and RNAi was performed as previously described (Goshima et al., 2007; Bettencourt-Dias and Goshima, 2009). Schneider's medium (Gibco) supplemented with 10% serum and antibiotics was used for cell culture. For imaging, cells were plated onto a 96-well, glass-bottom plate that was precoated with 5 μ l of 0.5 mg/ml concanavalin A solution (1 h at 24°C). For Klp61F RNAi, cells were treated with double-stranded RNA (dsRNA) targeting the 5' UTR region for 72 h and then plated. The primer sequences for dsRNA were 5'-TATTTGCGCATTATTTA AAATTG-3' and 5'-CATATTGATCAATTGAAAC-3' (Goshima and Vale, 2005), to which the T7 promoter sequences (5'-TAATACGAC TCACTATAGGG-3') were added. For in vitro transcription, the PCR product was mixed with T7 RNA polymerase and 7.5 mM ribonucleotide triphosphates and incubated at 37°C overnight (\sim 16 h). To complete annealing of dsRNA, the reaction mixture was incubated at 95°C for 5 min using a heat block and then cooled down gradually by leaving the block at room temperature. S2 cells in a 24-well plastic culture plate were treated with 5 μ g dsRNA in serum-free medium for 50 min, followed by serum addition. Live imaging of EB1-GFP was performed using spinning-disc confocal microscopy with a 100 \times 1.40 NA or 100 \times 1.45 NA objective lens (23–25°C). A CSU-X1 confocal unit (Yokogawa Electric Corporation) attached to a Nikon TE or Ti inverted microscope and EMCCD camera ImagEM (Hamamatsu Photonics) were used for image acquisition. The microscopes and attached devices were controlled using Micromanager. To eliminate actin cytoskeleton that might have physically prevented MT growth (Li et al., 2011), an actin inhibitor (2.5 μ M latrunculin A) was added to cells before imaging. In the Plk1^{Polo} inhibition experiment, cells were treated with control DMSO or 1 μ M BI 2536, which was shown to be effective in S2 cells (Januschke et al., 2013; Kachaner et al., 2014), and images were acquired at 3-s intervals.

Phos-tag SDS-PAGE, autoradiography, and immunoblotting

Phosphorylation of purified proteins by Plk1^{Polo}, except CLASP^{Mast/Orbit}, was evaluated by Phos-tag SDS-PAGE (Kinoshita et al., 2006). 1 mM ATP was added to the reaction. EB1 was run on an 8% acrylamide gel, whereas a 5% gel was used for tubulin, Sentin, and kinesin-13^{Plk10A}. The gel for XMAP215^{Msp}s was a mixture of 3% acrylamide and 0.5% agarose, wherein agarose assisted in gel polymerization. Acrylamide conjugated with the Phos-tag molecule at 20 μ M. For unknown reasons, the GFP-CLASP^{Mast/Orbit} band was not clearly detectable on any type of gel. CLASP^{Mast/Orbit} phosphorylation was therefore monitored by autoradiography, where 113 nM Plk1^{Polo} (wild type or KD) was mixed with 1 μ M GFP-CLASP^{Mast/Orbit} in MRB80/DTT with 100 mM KCl, 0.1 mM ATP (containing 110 nM ATP-P³²), and 80 mM β -glycerophosphate for 30 min at 24–25°C, followed by SDS-PAGE and Coomassie staining. Autoradiography was used to test for phosphorylation by Cdk1, where 27 nM recombinant Cdk1–cyclin B was mixed with 2.1 μ M EB1, 1.6 μ M tubulin, 1.7 μ M Sentin, 1.8 μ M kinesin-13^{Plk10A}, 1.2 μ M CLASP^{Mast/Orbit}, or 2.2 μ M XMAP215^{Msp}s. Immunoblotting of the S2 extract was performed with the following antibodies: anti- α -tubulin (DM1A, 1:1,000, mouse; Sigma-Aldrich), anti-CLASP^{Mast/Orbit} (1:300, rabbit; gift from H. Maiato, University of Porto, Porto, Portugal), anti-EB1 (1:1,000, rabbit; gift from S. Rogers, University of North Carolina, Chapel Hill, NC), anti-Klp10A (1:300, rabbit; gift from D. Sharp, Albert Einstein College of Medicine, Bronx, NY), anti-Sentin (1:1,000, rabbit; Li et al., 2011), and anti-XMAP215^{Msp}s (1:500, rabbit; gift from H. Ohkura, University of Edinburgh, Edinburgh, Scotland, UK).

Online supplemental material

Fig. S1 shows recombinant proteins used in this study, kinesin-13^{Kip10A} activity and a comparison between S2-derived tubulin and pig brain tubulin. Fig. S2 shows comparison of in vitro and in vivo parameters associated with MT dynamics. Fig. S3 compares the in vitro and in vivo effects of CLASP^{Mast/Orbit} and Sentin. Fig. S4 displays the effect of Cdk1–cyclin B in MT dynamics in vitro. In Fig. S5, cellular protein concentration of five factors and tubulin were estimated. Kinetic parameters of in vitro reconstituted MT polymerization dynamics are presented in Tables S1, S2, and S3.

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