

Synergistic role of fission yeast Alp16^{GCP6} and Mzt1^{MOZART1} in γ -tubulin complex recruitment to mitotic spindle pole bodies and spindle assembly

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ABSTRACT In fission yeast, γ -tubulin ring complex (γ TuRC)-specific components Gfh1^{GCP4}, Mod21^{GCP5}, and Alp16^{GCP6} are nonessential for cell growth. Of these deletion mutants, only *alp16 Δ* shows synthetic lethality with temperature-sensitive mutants of Mzt1^{MOZART1}, a component of the γ TuRC required for recruitment of the complex to microtubule-organizing centers. γ -Tubulin small complex levels at mitotic spindle pole bodies (SPBs, the centrosome equivalent in fungi) and microtubule levels for preanaphase spindles are significantly reduced in *alp16 Δ* cells but not in *gfh1 Δ* or *mod21 Δ* cells. Furthermore, *alp16 Δ* cells often form monopolar spindles and frequently lose a minichromosome when the spindle assembly checkpoint is inactivated. Alp16^{GCP6} promotes Mzt1-dependent γ TuRC recruitment to mitotic SPBs and enhances spindle microtubule assembly in a manner dependent on its expression levels. Gfh1^{GCP4} and Mod21^{GCP5} are not required for Alp16^{GCP6}-dependent γ TuRC recruitment. Mzt1 has an additional role in the activation of the γ TuRC for spindle microtubule assembly. The ratio of Mzt1 to γ TuRC levels for preanaphase spindles is higher than at other stages of the cell cycle. Mzt1 overproduction enhances spindle microtubule assembly without affecting γ TuRC levels at mitotic SPBs. We propose that Alp16^{GCP6} and Mzt1 act synergistically for efficient bipolar spindle assembly to ensure faithful chromosome segregation.

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INTRODUCTION

Microtubules play essential roles in many cellular processes, such as chromosome segregation, cell division, protein and organelle transport, cell polarity, and cell motility. It is crucial for the cell to spatially

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Abbreviations used: GFP, green fluorescent protein; γ TuC, γ -tubulin complex; γ TuRC, γ -tubulin ring complex; γ TuSC, γ -tubulin small complex; mCh, monomeric Cherry; mRFP, monomeric red fluorescent protein; MTOC, microtubule-organizing center; SPB, spindle pole body; YES, yeast extract and supplements.

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and temporally regulate microtubule assembly and its dynamics. γ -Tubulin complexes (γ TuCs) are essential for the initiation and regulation of microtubule assembly (reviewed in Kollman *et al.*, 2011; Teixidó-Travesa *et al.*, 2012; Lin *et al.*, 2015; Oakley *et al.*, 2015; Petry and Vale, 2015). γ TuCs localize to major microtubule-organizing centers (MTOCs), such as the centrosome in animal cells and the spindle pole body (SPB; the centrosome equivalent) in fungi, as well as to minor MTOCs, such as the nuclear envelope and the Golgi apparatus. In addition, in some systems, the γ TuCs localize along the length of microtubules to promote nucleation from the sides of other microtubules.

Two types of γ TuCs, the γ -tubulin small complex (γ TuSC) and the γ -tubulin ring complex (γ TuRC), have been identified in the cell (Kollman *et al.*, 2011; Lin *et al.*, 2015). The γ TuSC is composed of γ -tubulin, GCP2, and GCP3. The γ TuSC forms a γ TuRC ring structure in the presence of GCP4, GCP5, GCP6, and the small protein MOZART1 (Zheng *et al.*, 1995; Moritz *et al.*, 1995, 2000; Murphy *et al.*, 2001; Hutchins *et al.*, 2010). Down-regulation of GCP4, GCP5, or GCP6 disrupts γ TuRC assembly (Verollet *et al.*, 2006;

Izumi et al., 2008; Bahtz et al., 2012). The microtubule-nucleating activity of the γ TuRC is reminiscent of that of isolated centrosomes (Zheng et al., 1995) and much higher than that of the γ TuSC (Oegema et al., 1999).

Two models have been proposed for the roles of γ TuRC-specific components in the assembly of the γ TuRC ring. In one model, GCP4, GCP5, and GCP6 form a cap-like scaffold that arranges multiple γ TuSCs into a ring structure (Moritz et al., 2000; Zhang et al., 2000). In another, more recent model, GCP4, GCP5, and GCP6 are directly incorporated into the ring structure and bind to γ -tubulin (Kollman et al., 2011). The latter is supported by the observed sequence similarity among the GCPs, the structural similarity of GCP4 to GCP2 and GCP3 (Guillet et al., 2011), and the ability of *Drosophila* GCP5 and GCP6 to bind γ -tubulin (Gunawardane et al., 2003). In either case, interaction of the γ TuRC-specific components with the MTOCs seems to require their interacting with the γ TuSC, since they are not localized to the MTOCs without the γ TuSC (Xiong and Oakley, 2009).

In human cells, down-regulation of the γ TuRC-specific components GCP4, 5, or 6 reduces γ TuC localization at the spindle poles and induces monopolar spindle formation (Izumi et al., 2008; Bahtz et al., 2012). GCP4, 5, and 6, however, are nonessential in the fruit fly *Drosophila melanogaster*, fission yeast *Schizosaccharomyces pombe*, and filamentous fungus *Aspergillus nidulans* (Fujita et al., 2002; Schnorrer et al., 2002; Venkatram et al., 2004; Anders et al., 2006; Verollet et al., 2006; Xiong and Oakley, 2009). The budding yeast *Saccharomyces cerevisiae* does not possess MOZART1 or GCP4, 5, or 6 and only has the γ TuSC composed of Tub4/ γ -tubulin, Spc97/GCP2, and Spc98/GCP3. Recent studies of the *S. cerevisiae* γ TuSC showed that a ring structure could be assembled in vitro from the γ TuSC with the help of Spc110, a receptor of the γ TuSC at the nuclear side of the SPB (Kollman et al., 2010). Quantification of the number of γ TuSC components at microtubule nucleation sites in vivo supports the presence of a spiral of approximately seven γ TuSCs at the nucleation site (Erlemann et al., 2012). γ TuSC ring assembly may be restricted to SPBs in vivo and temporally regulated by the phosphorylation of the γ TuSC and Spc110 (Lin et al., 2014).

These observations imply that two types of ring structure may exist at MTOCs in higher eukaryotes and other fungi—the γ TuSC ring assembled on MTOCs from the γ TuSCs, and the γ TuRC ring recruited from the cytoplasm to the MTOCs. This also raises a question regarding the roles of MOZART1 and GCP4, 5, and 6 in microtubule assembly. In *S. cerevisiae*, the SPBs are the sole MTOCs and nucleate microtubules throughout the cell cycle, whereas most of the MTOCs in higher organisms and other fungi are transiently activated or assembled during the cell cycle. These differences in MTOC activities suggest that MOZART1 and γ TuRC-specific components are involved in spatial and temporal regulation of microtubule assembly. In accordance with this, it has been shown that the γ TuRC plays a role in microtubule-dependent microtubule nucleation (Verollet et al., 2006) and centriole biogenesis (Bahtz et al., 2012), and that MOZART1 acts as a recruitment factor for the γ TuRC to mitotic MTOCs in human cells (Hutchins et al., 2010; Teixidó-Travesa et al., 2010).

In fission yeast, any single deletion of the γ TuRC-specific components Gfh1^{GCP4}, Mod21^{GCP5}, and Alp16^{GCP6} and double and triple deletions are viable (Fujita et al., 2002; Venkatram et al., 2004; Anders et al., 2006). Single deletions show a reduction in the cytoplasmic microtubule number and interphase MTOC activity. Double and triple deletions do not lead to additional defects (Anders et al., 2006). No major mitotic defects have been reported in these deletions. In contrast, Mzt1^{MOZART1} is essential for cell growth and is required for the recruitment of γ TuCs to all the MTOCs (Masuda

et al., 2013; Dhani et al., 2013). We show here the role of Alp16^{GCP6} in mitotic spindle assembly. We find that Alp16^{GCP6} alone functions in mitosis for γ TuRC assembly without Gfh1^{GCP4} and Mod21^{GCP5} and that Alp16^{GCP6} and Mzt1 work together for efficient spindle microtubule assembly. We present a model for the synergistic effects of Alp16^{GCP6} and Mzt1 on mitotic γ TuRC assembly and activation.

RESULTS

***alp16 Δ* is synthetic lethal with a *mzt1* temperature-sensitive mutation**

We previously showed that Mzt1 is an essential γ TuRC component required for the recruitment of the complex to MTOCs (Masuda et al., 2013). To further study the role of Mzt1 in γ TuRC and γ TuSC function, we crossed *mzt1-T27W*—a temperature-sensitive *mzt1* mutant strain—with strains in which one of the genes encoding γ TuRC-specific components Gfh1^{GCP4}, Mod21^{GCP5}, or Alp16^{GCP6} is deleted. Tetrad dissection showed that *alp16 Δ* but not *gfh1 Δ* or *mod21 Δ* was synthetic lethal with the *mzt1-T27W* mutation (Figure 1A). We examined whether *alp16 Δ* shows synthetic lethality with temperature-sensitive mutants of the other γ TuSC components and found that *alp16 Δ* but not *gfh1 Δ* or *mod21 Δ* was synthetic lethal with *alp4-1891*, a temperature-sensitive mutant of Alp4^{GCP2} (Vardy and Toda, 2000; Supplemental Figure S1A).

Alp16^{GCP6} is localized to mitotic SPBs without Gfh1^{GCP4} or Mod21^{GCP5} and required for localization of Gfh1^{GCP4} and Mod21^{GCP5} to mitotic SPBs

We studied localization of Gfh1^{GCP4}, Mod21^{GCP5}, and Alp16^{GCP6} at mitotic SPBs using strains expressing the proteins tagged with N-terminal green fluorescent protein (GFP), Sid4—monomeric red fluorescent protein (mRFP), and monomeric Cherry (mCh)—Atb2 (Figure 1B). Sid4-mRFP and mCh-Atb2 were used as SPB and microtubule markers, respectively. GFP-Alp16^{GCP6} localized to the mitotic SPBs in both *gfh1 Δ* and *mod21 Δ* cells. In contrast, GFP-Gfh1^{GCP4} or GFP-Mod21^{GCP5} did not localize to the mitotic SPBs in *alp16 Δ* cells. Gfh1^{GCP4} needed Alp16^{GCP6} but not Mod21^{GCP5} for localization to the mitotic SPBs. Mod21^{GCP5} relied on both Alp16^{GCP6} and Gfh1^{GCP4} for mitotic SPB localization. These results show that the three γ TuRC-specific components constitute localization hierarchy: Alp16^{GCP6} acts most upstream, which is followed by Gfh1^{GCP4} and Mod21^{GCP5}. Despite the synthetic lethality with *mzt1* and *alp4* mutants and the disappearance of Gfh1^{GCP4} and Mod21^{GCP5} from mitotic SPBs, *alp16 Δ* did not cause an obvious growth reduction (Supplemental Figure S1, D and E).

Levels of γ TuC at mitotic SPBs and levels of spindle microtubules are significantly reduced in *alp16 Δ* cells

To study the effect of *alp16 Δ* on mitotic progression in detail, we observed the dynamic behavior of γ TuCs and spindle microtubules in wild-type and *alp16 Δ* cells expressing Sid4-mRFP, mCh-Atb2, and GFP-Alp4^{GCP2} or GFP-Mzt1 (Figure 2, A and B). We found that the levels of both Alp4^{GCP2} and Mzt1 at mitotic SPBs and the levels of spindle microtubules were reduced in *alp16 Δ* cells compared with wild-type cells. Quantitative analysis showed that GFP-Alp4^{GCP2} and GFP-Mzt1 levels at both mitotic and interphase SPBs in *alp16 Δ* cells were reduced to 20–40% of wild-type levels (Figure 2, C and D, and Supplemental Figure S2, C–I). Spindle microtubule levels in *alp16 Δ* cells were quantified using GFP-Atb2 to avoid any potential inhibitory effects of GFP tagging of Alp4^{GCP2} and Mzt1 on microtubule nucleation and prevent inaccurate measurement of microtubule levels due to fast photobleaching observed with mCh-Atb2. We found that spindle microtubule levels in *alp16 Δ* cells were reduced to

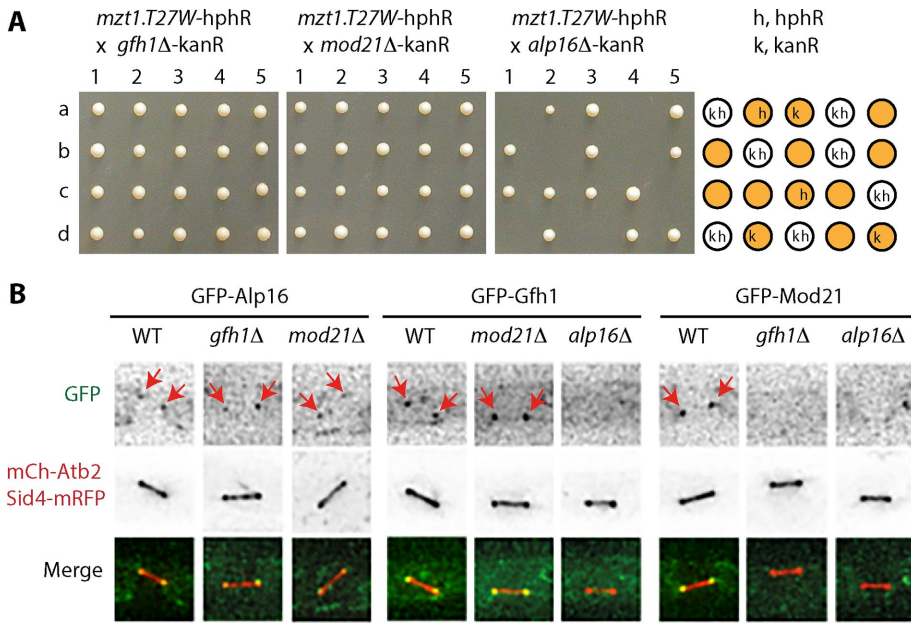


FIGURE 1: *alp16Δ* is synthetic lethal with the *mzt1-T27W* mutant, and Alp16^{GCP6} localizes to mitotic SPBs in the absence of Gfh1^{GCP4} and Mod21^{GCP5}. (A) *alp16Δ* is synthetic lethal with the *mzt1-T27W* mutant. Diploid cells obtained from crosses between *mzt1-T27W-hphR* and *gfh1Δ-kanR*, *mod21Δ-kanR*, or *alp16Δ-kanR* were sporulated, and individual spores (a–d) in each ascus (1–5) were dissected on YES plates. Nonviable spores from the cross between *mzt1-T27W* and *alp16Δ* germinated, divided several times, and arrested with elongated morphologies. Drug resistance of viable spores (orange circle) and predicted resistance of nonviable spores (white circle): k, *kanR* (resistance to G418); h, *hphR* (resistance to hygromycin). (B) Localization of Alp16^{GCP6}, Gfh1^{GCP4}, and Mod21^{GCP5} to mitotic SPBs. Localization of GFP-Alp16^{GCP6}, GFP-Gfh1^{GCP4}, and GFP-Mod21^{GCP5} to mitotic SPBs was examined in wild-type (WT), *gfh1Δ*, *mod21Δ*, and *alp16Δ* cells containing mCh-Atb2 and Sid4-mRFP. Scale bar, 10 μm.

50–60% of wild-type levels during early M phase and 70–80% during mid M phase (Figure 2E).

Alp16-dependent microtubule assembly is activated in preanaphase spindles

To determine which phase of mitosis is affected by *alp16Δ*, we plotted levels of Alp4^{GCP2}, Mzt1, and spindle microtubules against spindle length (Figure 2, C–E). In wild-type cells, levels of Alp4^{GCP2} and Mzt1 constantly increased, along with increasing spindle length, whereas spindle microtubule levels were high in spindles 1–3 μm long, and levels then decreased when spindles further elongated. Thus, spindle microtubule levels were high in preanaphase spindles, since anaphase spindle elongation was found to begin at a length of ~3 μm on average (Supplemental Figure S3). On the other hand, in *alp16Δ* cells, Alp4^{GCP2} and Mzt1 levels were ~30% of that of wild-type levels upon spindle assembly, which gradually increased as mitosis progressed (Figure 2, C and D). Spindle microtubule levels in *alp16Δ* cells were ~60% of wild-type levels upon spindle assembly, gradually increased until metaphase, and then decreased as the spindle further elongated at anaphase (Figure 2E). The ratio of Alp4^{GCP2} and Mzt1 levels in *alp16Δ* cells to wild-type cells did not change much during mitosis. In contrast, the ratio of spindle microtubule levels in *alp16Δ* cells to wild-type cells was ~60% at prometaphase through metaphase, which then increased to 70% in early anaphase and further increased to 90–100% in late anaphase (Figure 2E). These results suggest that Alp16^{GCP6}-dependent spindle microtubule assembly is prominent during early M phase, especially in preanaphase spindles.

To study whether the deletion of *gfh1* or *mod21*—two genes encoding other γTuRC-specific components—results in the phenotypes similar to *alp16Δ*, we measured the levels of γTuC at the mitotic SPBs and of spindle microtubules from preanaphase spindles (<3.5 μm in length; Figure 2F). Alp4^{GCP2} levels in *gfh1Δ* and *mod21Δ* cells were instead increased to 147 and 111% compared with wild-type cells, whereas Mzt1 levels in *gfh1Δ* and *mod21Δ* cells were decreased to 88 and 72%, respectively. Spindle microtubule levels decreased to 79 and 75% in *gfh1Δ* and *mod21Δ* cells compared with wild-type cells. In contrast, in *alp16Δ* cells, levels of Alp4, Mzt1, and spindle microtubules were reduced to 46, 33, and 39% of wild-type levels, respectively. Thus the *alp16Δ* deletion more specifically and significantly reduced the levels of γTuC at the mitotic SPBs and of spindle microtubules than did the deletion of *gfh1* or *mod21*.

Monopolar spindles are frequently observed before bipolar spindle assembly in *alp16Δ* cells

Our results suggest that the number of microtubules assembled from mitotic SPBs in *alp16Δ* cells is reduced to 60% compared with wild-type cells. To explore the possibility that bipolar spindle assembly is temporarily affected in *alp16Δ* cells, we observed spindle assembly in *alp16Δ* cells expressing GFP-Alp4^{GCP2} and mCh-Atb2 without Sid4-mRFP labeling, which hindered observation of spindle microtubule behavior. About 90% of *alp16Δ* cells showed monopolar spindles or bipolar spindles with extra microtubules emanating from one or both SPBs for 1–3 min, which eventually transformed into stable bipolar spindles (Figure 3, A and B). In contrast, ~60% of wild-type cells showed monopolar or bipolar spindles with extra microtubules for ~1 min, before stable, thick bipolar spindles were assembled (Figure 3B).

In addition to transient monopolar spindle assembly, *alp16Δ* cells remained for longer at the postmitotic stage with equatorial MTOCs, where Alp4^{GCP2} and Mzt1 levels were increased twofold to threefold over wild-type levels (Supplemental Figure S4, A–C). Alp4^{GCP2} and Mzt1 in interphase cells tended to form cytoplasmic dots besides interphase MTOCs, some of which did not seem to be involved in cytoplasmic microtubule formation (Supplemental Figure S4D).

Minichromosome loss rate of *alp16Δ* cells is higher than that of wild-type cells and significantly increased with inactivation of spindle assembly checkpoint

alp16Δ cells may show a defect in chromosome segregation if the chromosome number is artificially increased, since reduction in the number of spindle microtubules seems to mostly affect the formation of kinetochore microtubules. To test this possibility, we introduced a minichromosome Ch16 carrying the *ade6-M216* mutation (Niwa et al., 1986) in *alp16Δ* cells that contained the *ade6-M210* mutation. Loss of the minichromosome was detected as red colony or red-sector colony formation (see Materials and Methods). We

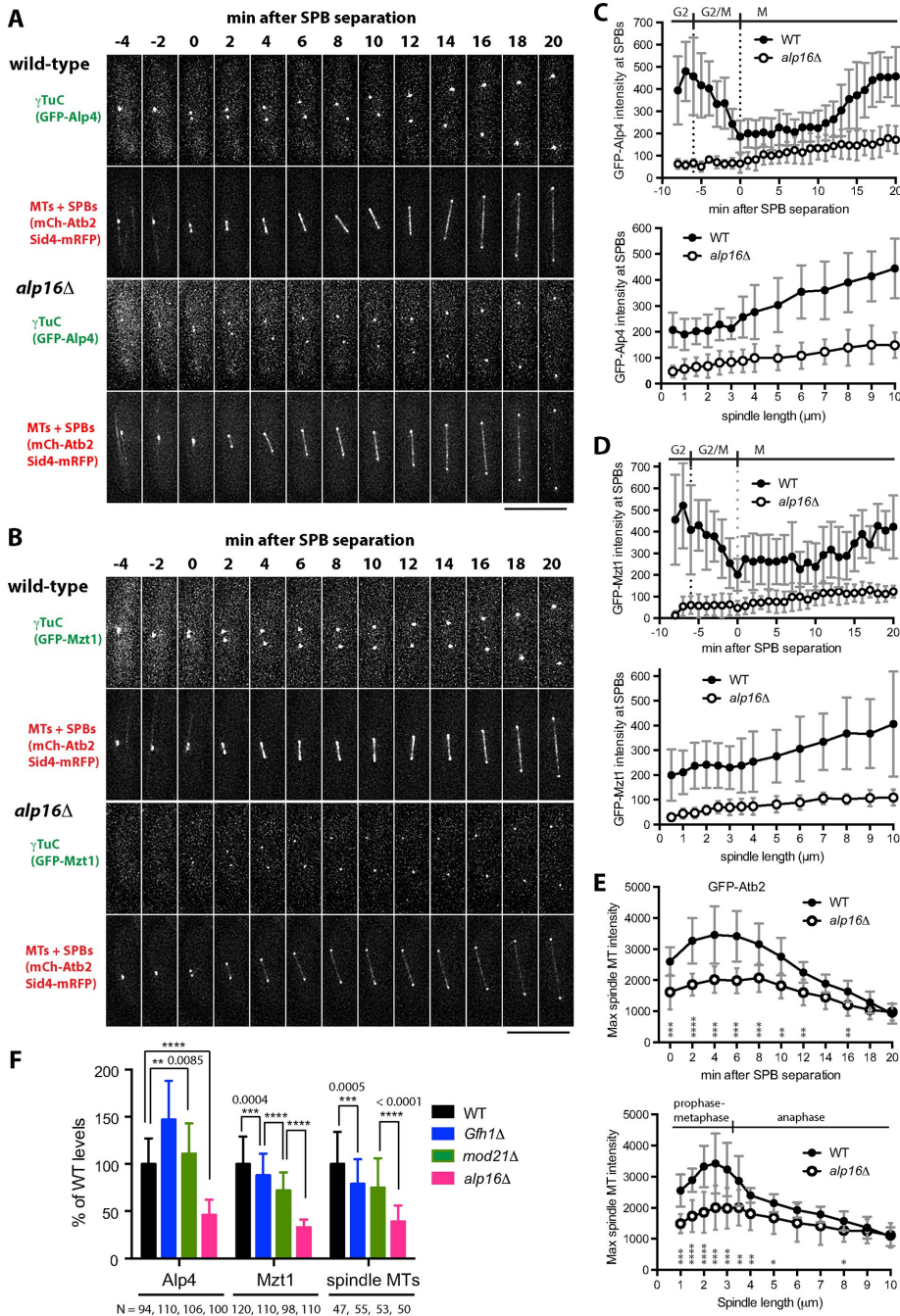


FIGURE 2: *alp16Δ* reduces levels of γ TuC at mitotic SPBs and of spindle microtubule levels at early M phase. (A, B) γ TuC levels at mitotic SPBs and spindle MT levels are reduced in *alp16Δ* cells. Time-lapse images of wild-type and *alp16Δ* cells containing mCh-Atb2 (for MTs), Sid4-mRFP (for SPBs), and either GFP-Alp4^{GCP2} (A) or GFP-Mzt1 (B) (for γ TuC). Time 0 is the time when the two SPBs are separated for spindle assembly. Scale bar, 10 μ m. (C, D) Quantification of γ TuC levels. Average intensity of GFP signal at mitotic SPBs in wild-type and *alp16Δ* cells expressing GFP-Alp4^{GCP2} (C; $N = 9$ and 8 , respectively) or GFP-Mzt1 (D; $N = 12$ and 9 , respectively) as a function of time after SPB separation or spindle length. SDs are shown as gray bars. (E) Quantification of spindle microtubule levels. Average intensity of GFP signal for spindles from wild-type ($N = 10$) and *alp16Δ* ($N = 12$) cells expressing GFP-Atb2 as a function of time after SPB separation or spindle length. SDs are shown as gray bars. The p values from unpaired t test are shown at each time point or spindle length. * $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$, and **** $p < 0.0001$. (F) Comparison of γ TuC levels at the mitotic SPBs and spindle microtubule levels in wild-type, *gfh1Δ*, *mod21Δ*, and *alp16Δ* cells. γ TuC levels at SPBs and spindle microtubule levels from spindles $< 3.5 \mu$ m in length were measured in wild-type, *gfh1Δ*, *mod21Δ*, and *alp16Δ* cells containing GFP-Alp4^{GCP2} or GFP-Mzt1 and mCh-Atb2. The p values from unpaired t test are shown.

found that *alp16Δ* cells showed a higher loss rate than wild-type cells. Furthermore, *alp16Δ mad2Δ* cells showed much higher loss rates, with the spindle assembly checkpoint inactivated by *mad2* deletion (Figure 3C). These results indicate that Alp16^{GCP6}-dependent microtubule nucleation ensures faithful chromosome segregation.

Artificial targeting of Alp16^{GCP6} to Pcp1 at SPBs decreases γ TuC levels but increases the ratio of spindle microtubules to γ TuC levels

Our results suggest that Alp16^{GCP6} is involved in the recruitment of the γ TuRC to mitotic SPBs. Although γ TuRC assembly seems to be a prerequisite for localization of γ TuRC-specific components to the MTOCs, it is not known whether artificial targeting of the components to the MTOCs affects γ TuC levels. We tested whether targeting Alp16^{GCP6} to mitotic SPBs increases γ TuRC levels at the SPBs and spindle microtubule levels. Because Pcp1 is likely to act as a receptor for the γ TuRC at the mitotic SPB (Fong *et al.*, 2010), we tagged Pcp1 with the GFP-binding protein (GBP; Rothbauer *et al.*, 2008) and used it to target GFP-Alp16^{GCP6} to mitotic SPBs (Figure 4, A and B). GFP-Alp16^{GCP6} levels at mitotic SPBs and spindle microtubule levels were increased in cells expressing both GFP-Alp16^{GCP6} and GBP-Pcp1 (Figure 4, A, C, and E). γ TuC levels, represented by mCh-Alp4, at mitotic SPBs, however, decreased in the cells with artificial targeting of Alp16^{GCP6} to Pcp1 (Figure 4, B, D, and E). Thus targeting of Alp16^{GCP6} to Pcp1 had an inhibitory effect on recruitment of the γ TuRC or γ TuSC to the SPBs. This reduction is probably due to occupation of the γ TuC-binding site on Pcp1. Nonetheless, spindle microtubule levels and the ratio of spindle microtubules to γ TuC levels at the mitotic SPBs were increased (Figure 4E). These results imply that the γ TuRC ring recruited to the SPBs has higher microtubule nucleation activity or assembles more stable microtubules than the nucleating complex assembled from γ TuSCs at the SPBs.

Increasing expression levels of Alp16^{GCP6} results in increased levels of γ TuC at mitotic SPBs and of spindle microtubules independently of Gfh1^{GCP4} and Mod21^{GCP5}

To further investigate whether the increase in levels of Alp16^{GCP6} at the mitotic SPBs affects γ TuC levels and spindle microtubule levels, we changed the expression levels of Alp16^{GCP6}. The *alp16* gene was integrated at the *lys1* locus in *alp16Δ* cells, and expression levels were controlled

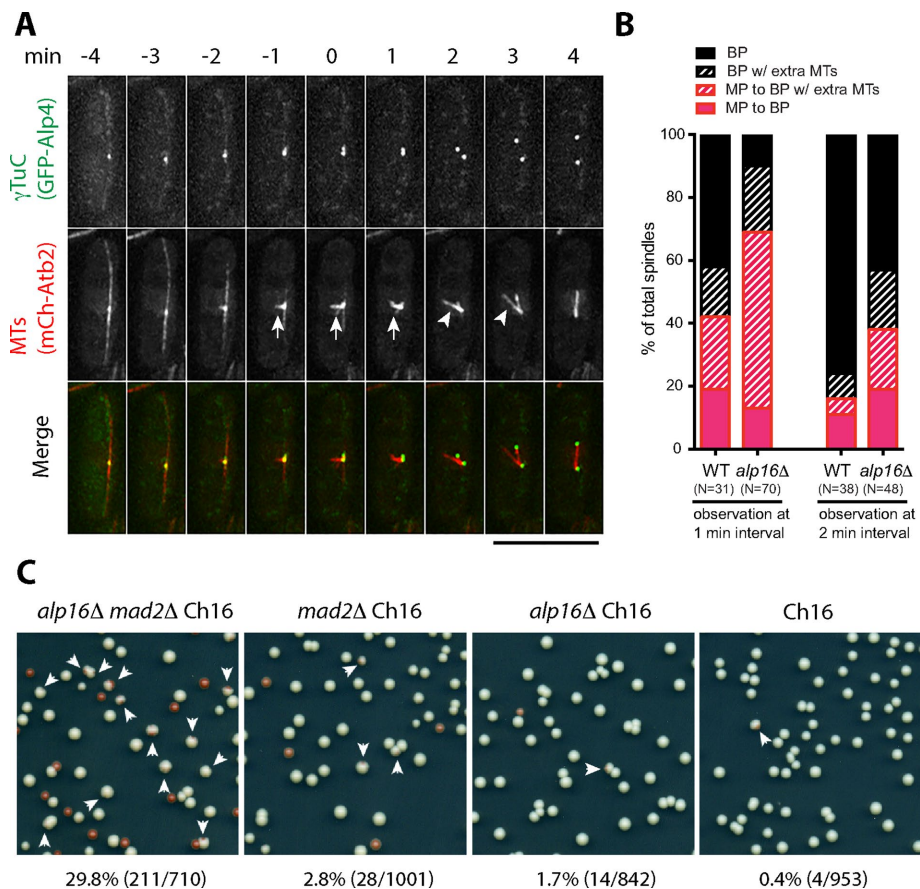


FIGURE 3: *alp16Δ* affects spindle assembly and chromosome segregation. (A) Monopolar spindles observed during early M phase in *alp16Δ* cells. Time-lapse images of an *alp16Δ* cell containing GFP-Alp4^{GCP2} and mCh-Atb2 show formation of a monopolar spindle (arrow), which eventually transforms into a bipolar spindle. A bipolar spindle with an extra microtubule bundle nucleating from the SPB (arrowhead) is observed at 2 and 3 min after SPB separation. Scale bar, 10 μ m. (B) Percentages of wild-type and *alp16Δ* mitotic cells with monopolar (MP) spindles or bipolar (BP) spindles with extra microtubule bundles from time-lapse imaging at 1- and 2-min intervals. MP to BP, monopolar spindles transform into bipolar spindles. MP to BP w/extra MTs, monopolar spindles transform into bipolar spindles with extra microtubule bundles, which eventually transform into bipolar spindles. BP w/extra MTs, without monopolar spindle assembly, bipolar spindles with extra microtubule bundles form, which eventually transform into bipolar spindles. BP, bipolar spindles form without monopolar spindle assembly. (C) *alp16Δ* cells frequently lose a minichromosome in the absence of Mad2. *alp16Δ mad2Δ ade6-M210* Ch16, *mad2Δ ade6-M210* Ch16, *alp16Δ ade6-M210* Ch16, and *ade6-M210* Ch16 cells were grown at 27°C on YE plates for 6 d. Arrows indicate red-sectored colonies that lost Ch16 during the 6 d of growth. The chromosome loss rate was calculated as the percentage of red-sectored colonies out of the sum of red-sectored and white colonies.

from a thiamine-repressible *nmt1* promoter and its derivatives that have lower activities (Basi *et al.*, 1993). We found that the levels of γ TuC at the SPBs and levels of spindle microtubules increased along with the increase in expression levels of Alp16^{GCP6} (Figure 5, A–C).

To study the effect of Alp16^{GCP6} expression on γ TuC levels and spindle microtubule levels in more detail, we plotted spindle microtubule levels against γ TuC levels (Figure 5D, left). We then plotted ratio of spindle microtubules to γ TuC levels at the mitotic SPBs relative to those of *alp16Δ* and wild-type cells against γ TuC levels (Figure 5D, right). At a low expression level (under the control of P81*nmt* promoter in the presence of thiamine), γ TuC levels at mitotic SPBs were not increased significantly compared with levels in *alp16Δ* cells (by ~11% of wild-type levels). However, spindle

microtubule levels significantly increased by ~27% compared with those in *alp16Δ* cells. The ratio of spindle microtubules to γ TuC levels increased by ~29% compared with *alp16Δ* cells with similar γ TuC levels, suggesting again that the γ TuRC ring has a higher activity or forms more stable microtubules compared with the nucleating complex assembled from γ TuSCs without Alp16. At medium expression levels (under the control of P41*nmt* and P3*nmt* promoters in the presence of thiamine), spindle microtubule levels reached those of wild-type cells (~97%). with lower levels of γ TuC at the SPBs (~77%), or increased to levels higher than wild type (~118%), with levels of γ TuC similar to those of wild-type cells (~91%). At high expression levels (with P81 and P41*nmt* promoters in the absence of thiamine), both γ TuC levels at SPBs and spindle microtubule levels were higher than those in wild-type cells (Figure 5, B and D). At these medium and high expression levels, the ratio of spindle microtubules to γ TuC levels increased compared with wild-type cells.

To study whether Gfh1^{GCP4} and Mod21^{GCP5} are required for the Alp16^{GCP6}-dependent increase in levels of γ TuC at mitotic SPBs and of spindle microtubules, we examined localization of GFP-Gfh1^{GCP4} and GFP-Mod21^{GCP5} in cells overproducing Alp16^{GCP6}. Both components were found to localize at the mitotic SPBs (Figure 5E). Next, we overexpressed Alp16^{GCP6} in *alp16Δ gfh1Δ* and *alp16Δ gfh1Δ mod21Δ* cells and found that the Alp16^{GCP6}-dependent increase in levels of γ TuC at the mitotic SPBs and of spindle microtubules did not require Gfh1^{GCP4} or Mod21^{GCP5} (Figure 5F).

Alp16^{GCP6} overproduction suppresses the temperature sensitivity of *mzt1* mutant cells but not the lethality of *mzt1Δ* cells

Our results suggest a role for Alp16^{GCP6} in the recruitment of the γ TuRC to SPBs. Be-

cause Mzt1 is an essential component that acts as a recruitment factor for the γ TuRC (Masuda *et al.*, 2013), we examined whether increased levels of Alp16^{GCP6} suppress the phenotype of *mzt1* temperature-sensitive mutants. Serial dilution spot assays showed that overproduction of Alp16^{GCP6} suppressed the temperature sensitivity of *mzt1-T27W* (Figure 6A). Next, to test whether Alp16^{GCP6} overproduction rescues the lethality of the *mzt1* deletion, we constructed diploid cells in which Alp16^{GCP6} was overproduced and one of the *mzt1* genes was deleted. Tetrad analysis showed that Mzt1 was essential for growth even in the cells overproducing Alp16^{GCP6} (Figure 6B). This result suggests that Alp16^{GCP6}, when overproduced, could rescue compromised Mzt1 functions but still is not capable of substituting for its essential roles.

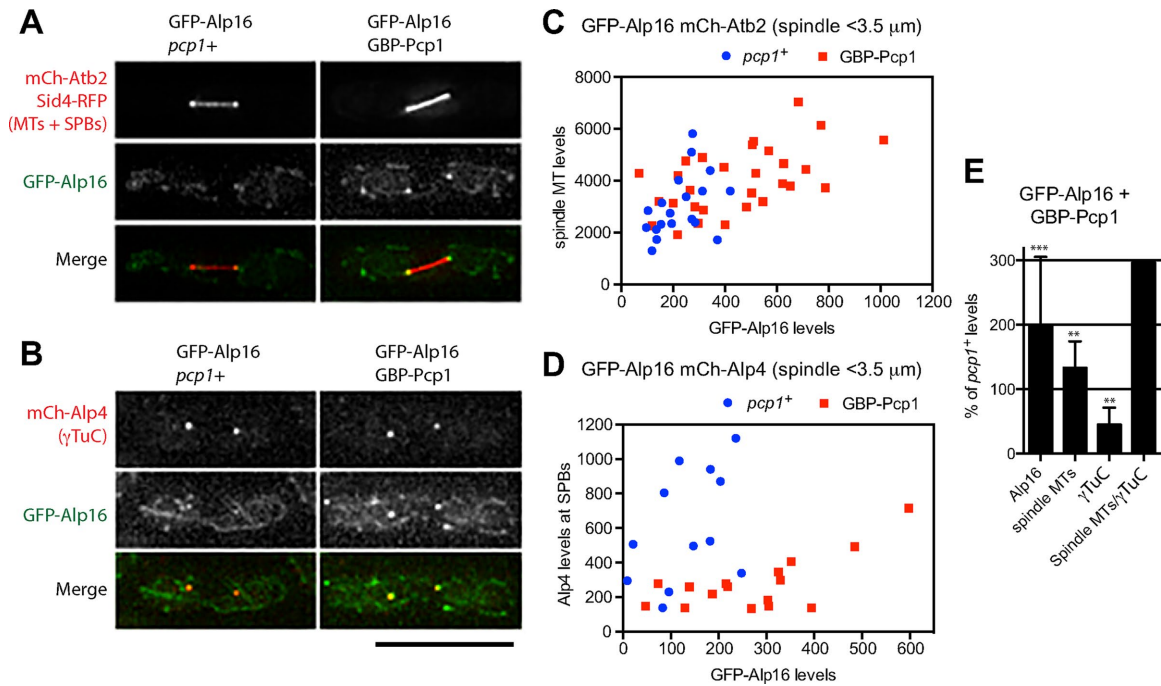


FIGURE 4: Artificial targeting of Alp16^{GCP6} to Pcp1 increases ratio of spindle microtubules to γ TuC levels. (A) Targeting of GFP-Alp16^{GCP6} to GBP-Pcp1 in cells expressing mCh-Atb2 increases spindle microtubule levels. Spindles observed in GFP-Alp16^{GCP6} and GFP-Alp16^{GCP6} GBP-Pcp1 cells expressing mCh-Atb2 and Sid4-mRFP. (B) Targeting of GFP-Alp16^{GCP6} to GBP-Pcp1 in cells expressing mCh-Alp4^{GCP2} reduces γ TuC levels at mitotic SPBs. Spindles observed in GFP-Alp16^{GCP6} and GFP-Alp16^{GCP6} GBP-Pcp1 cells expressing mCh-Alp4^{GCP2}. Scale bar, 10 μ m. (C) Quantification of spindle microtubule levels at mitotic SPBs in cells with or without Alp16^{GCP6} targeting to Pcp1. GFP-Alp16^{GCP6} and mCh-Atb2 levels for spindles <3.5 μ m in length were quantified. The sum of GFP-Alp16 levels at two SPBs of mitotic spindles is plotted against mCh-Atb2 levels. (D) Quantification of γ TuC levels at mitotic SPBs in cells with or without Alp16^{GCP6} targeting to Pcp1. GFP-Alp16^{GCP6} and mCh-Alp4^{GCP2} levels for spindles <3.5 μ m in length were quantified. (E) Levels of Alp16^{GCP6}, spindle microtubules, and γ TuC, and the ratio of spindle MTs to γ TuC levels in GFP-Alp16^{GCP6} GBP-Pcp1 cells were compared with those in GFP-Alp16^{GCP6} *pcp1+* cells. The *p* value is from unpaired *t* test: ****p* = 0.0003 for Alp16^{GCP6} (*N* = 19 for *pcp1+* and 29 for GBP-Pcp1 cells), ***p* = 0.0075 for spindle microtubules (*N* = 19 for *pcp1+* and 29 for GBP-Pcp1 cells), and ***p* = 0.0017 for γ TuC (*N* = 12 for *pcp1+* and 16 for GBP-Pcp1 cells).

Mzt1 increases spindle microtubule levels without affecting γ TuC levels at mitotic SPBs in wild-type cells

Because augmented Alp16^{GCP6} suppressed the temperature sensitivity of *mzt1* mutants, we examined whether higher levels of Mzt1 in turn suppress the phenotypes of *alp16 Δ* cells. Mzt1 overproduction in *alp16 Δ* cells had no effect on γ TuC levels at mitotic SPBs or spindle microtubule levels (Figure 7A). Unexpectedly, however, overproduction of Mzt1 in wild-type cells increased spindle microtubule levels without affecting γ TuC levels at mitotic SPBs (Figure 7A). To determine whether the ratio of Mzt1 to Alp4^{GCP2} is constant or variable during mitotic progression, we calculated the ratio of GFP-Mzt1 to GFP-Alp4^{GCP2} using the average intensity of GFP signal obtained from mitotic GFP-Mzt1 and GFP-Alp4^{GCP2} cells (Figure 2, C and D, and Supplemental Figure S5) and plotted it against spindle length (Figure 7B). The ratio was found to be high in preanaphase spindles 1–3 μ m in length compared with that of anaphase spindles in wild-type cells. The ratio did not change substantially in *alp16 Δ* cells and stayed low compared with wild-type cells. To confirm the increase in ratio of Mzt1 to Alp4^{GCP2} at early M phase, we directly measured Mzt1 and Alp4^{GCP2} levels in GFP-Alp4^{GCP2} mCh-Mzt1 cells. The ratio of Mzt1 to Alp4^{GCP2} in mitotic cells with preanaphase spindles was found to be higher than that in interphase cells (Figure 7C), suggesting that the number of Mzt1 proteins per γ TuRC increases at early M phase.

DISCUSSION

Roles of γ TuRC-specific components in spindle microtubule assembly

We showed here that Alp16^{GCP6} but not Gfh1^{GCP4} or Mod21^{GCP5} has demonstrable mitotic function in fission yeast. Levels of γ TuC at mitotic SPBs and spindle microtubule levels are significantly reduced in *alp16 Δ* cells. This reduction is not observed in *gfh1 Δ* or *mod21 Δ* cells. We also found that Alp16^{GCP6} expression in *alp16 Δ* and *alp16 Δ gfh1 Δ mod21 Δ* cells results in an increase in these levels (Figure 5). These Alp16^{GCP6}-specific effects are consistent with the interaction hierarchy of the γ TuRC-specific components with the γ TuSC (Anders *et al.*, 2006) and the hierarchy of localization of components to mitotic SPBs observed in this study: Alp16^{GCP6} alone can interact with the γ TuSC and localize to mitotic SPBs without Gfh1^{GCP4} and Mod21^{GCP5}, whereas Gfh1^{GCP4} and Mod21^{GCP5} require Alp16^{GCP6} for interaction with the γ TuSC and for localization to mitotic SPBs. Without Alp16^{GCP6}, the number of spindle microtubules nucleated from the SPBs is reduced but is still large enough to form kinetochore microtubules for all three chromosomes in fission yeast without any noticeable mitotic delay. Monopolar spindle assembly is observed more frequently but is eventually converted into a bipolar spindle. Increasing the number of chromosomes from three to four by introduction of a minichromosome reduces the number of microtubules used for kinetochore microtubule formation and dramatically

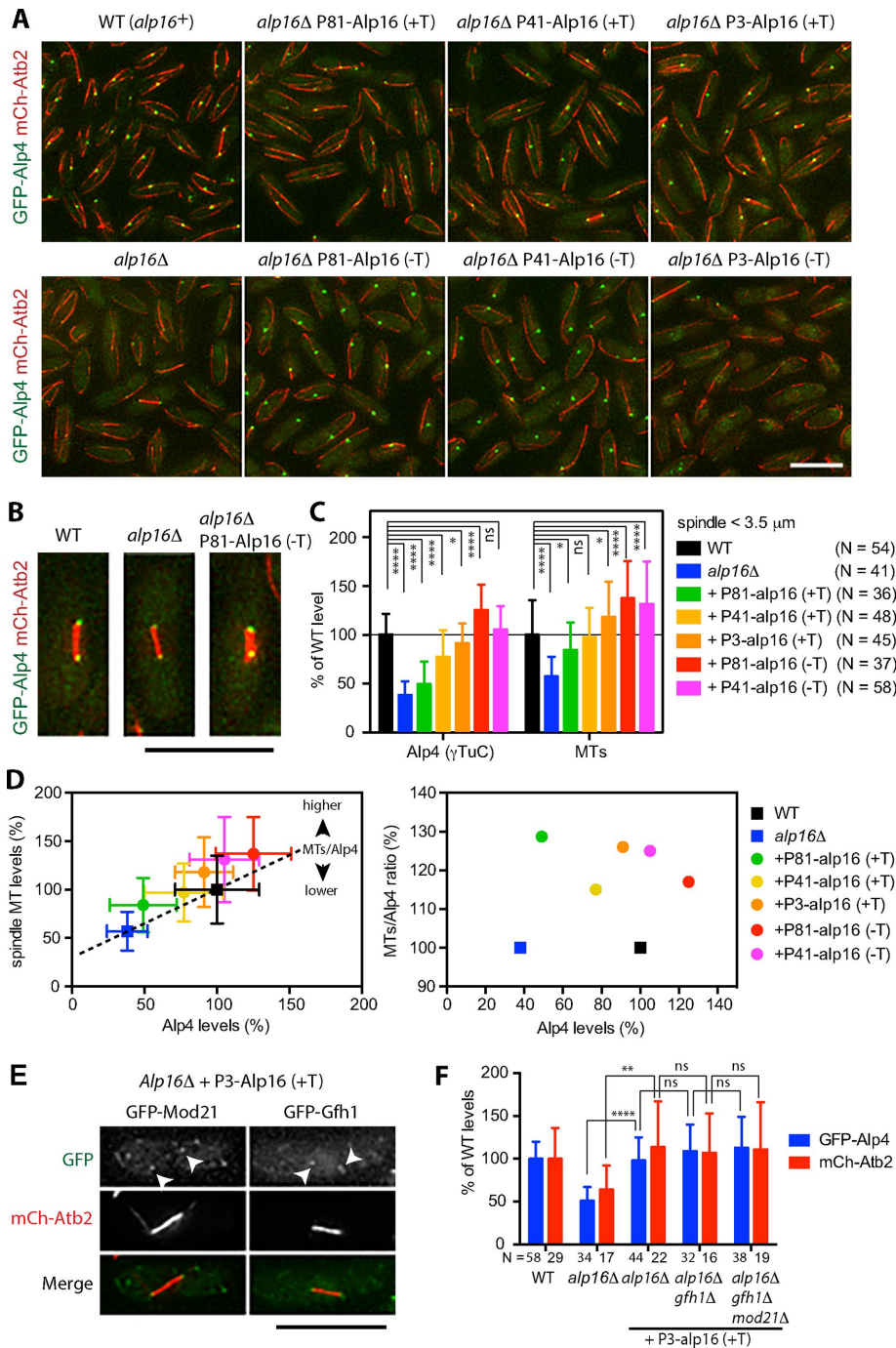


FIGURE 5: Elevated $Alp16^{GCP6}$ expression increases γ TuC levels at mitotic SPBs and spindle microtubule levels. (A) Increase in expression levels of $Alp16^{GCP6}$ increases γ TuC levels at SPBs. $Alp16^{GCP6}$ was expressed at the *lys1* locus in $alp16\Delta$ cells containing GFP- $Alp4^{GCP2}$ and mCh-Atb2 under the control of thiamine-repressible P81*nmt*, P41*nmt*, or P3*nmt* promoter in the presence (+T) and absence (-T) of 2 μ M thiamine at 27°C for 22–24 h. $Alp16^{GCP6}$ expression levels were the lowest under the P81*nmt* promoter with thiamine (P81-Alp16 (+T)) and increased in the following order: P81 (+T) < P41 (+T) < P3 (+T) < P81 (-T) < P41 (-T) < P3 (-T). The intensity of GFP- $Alp4^{GCP2}$ signals at SPBs increases along with the increase in expression levels of $Alp16^{GCP6}$ but decreases at the highest expression with P3*nmt* promoter in the absence of thiamine. Scale bar, 10 μ m. (B) Typical images of spindles observed in wild-type, $alp16\Delta$, and $alp16\Delta$ cells with P81*nmt*-Alp16 (-T). Note that spindles look thicker with higher levels of γ TuC at SPBs. (C) Elevated $Alp16^{GCP6}$ expression increases γ TuC levels at mitotic SPBs and spindle microtubule levels. Quantification of $Alp4^{GCP2}$ (γ TuC) and microtubule levels for spindles <3.5 μ m in length. The *p* values are from unpaired *t* test: *****p* < 0.0001, **p* < 0.05, and ns, *p* > 0.05. (D) The ratio of spindle microtubule to γ TuC level increases when $Alp16^{GCP6}$ is expressed. Spindle microtubule level was plotted against $Alp4^{GCP2}$ level using the data used for

increases the minichromosome loss rate when the spindle assembly checkpoint is inactivated. Thus $Alp16^{GCP6}$ -dependent spindle microtubule assembly ensures faithful chromosome segregation.

GCP6-specific phenotypes of mitotic spindle functions have been observed in other organisms. In *Aspergillus nidulans*, GCPD/GCP4, GCPE/GCP5, and GCPF/GCP6 exhibit a hierarchy of localization to the SPB similar to our results (Xiong and Oakley, 2009). *GCP6* Δ diploid cells lose chromosomes at a higher frequency, and *GCP6* Δ *mad2* Δ diploid cells show more severe synthetic sickness (Xiong and Oakley, 2009). In human cells, down-regulating GCP6 induces a high percentage of monopolar spindles compared with down-regulating GCP4 and GCP5 (Bahtz et al., 2012). These observations suggest that GCP4-6 mitotic functions are substantially conserved in the phylogenetically distant fungi *S. pombe* and *A. nidulans* and partly conserved in fungi and higher eukaryotes.

In contrast to the mitotic phenotypes, a single disruptant of *gfh1*, *mod21*, or *alp16* and double and triple disruptants reduce cytoplasmic microtubule number and interphase MTOC activity to a similar extent (Fujita et al., 2002; Anders et al., 2006), suggesting that the deletion of any γ TuRC-specific component disrupts interphase γ TuRC function. Differences in the properties of mitotic and interphase γ TuRCs may arise from the cell cycle-dependent

C. The dotted line was drawn connecting $alp16\Delta$ and wild-type levels, assuming that the increase in $Alp4^{GCP2}$ level increases spindle microtubule level linearly ($y = 0.69x + 0.31$, where $x = Alp4^{GCP2}$ level and $y =$ spindle microtubule level). $Alp16^{GCP6}$ expression increases spindle microtubule level to higher than the expected value from the equation. The ratio of spindle microtubule to $Alp4^{GCP2}$ level was calculated as a percentage of the expected value and plotted against the $Alp4^{GCP2}$ level. (E) $Gfh1^{GCP4}$ and $Mod21^{GCP5}$ localize to mitotic SPBs when $Alp16^{GCP6}$ is overexpressed in $alp16\Delta$ cells. Localization of GFP- $Gfh1^{GCP4}$ and GFP- $Mod21^{GCP5}$ was observed in $alp16\Delta$ cells with P3*nmt*-Alp16 (+T). Scale bar, 10 μ m. (F) $Gfh1^{GCP4}$ and $Mod21^{GCP5}$ are not required for $Alp16^{GCP6}$ -dependent γ TuC recruitment and spindle microtubule assembly. $Alp4^{GCP2}$ (γ TuC) and microtubule levels for spindles <3.5 μ m in length were quantified in wild-type, $alp16\Delta$, $alp16\Delta$ with P3-Alp16 (+T), $alp16\Delta$ *gfh1* Δ with P3-Alp16 (+T), and $alp16\Delta$ *gfh1* Δ *mod21* Δ cells with P3-Alp16 (+T). The *p* values are from unpaired *t* test: *****p* < 0.0001, ***p* < 0.01, and ns, *p* > 0.05.

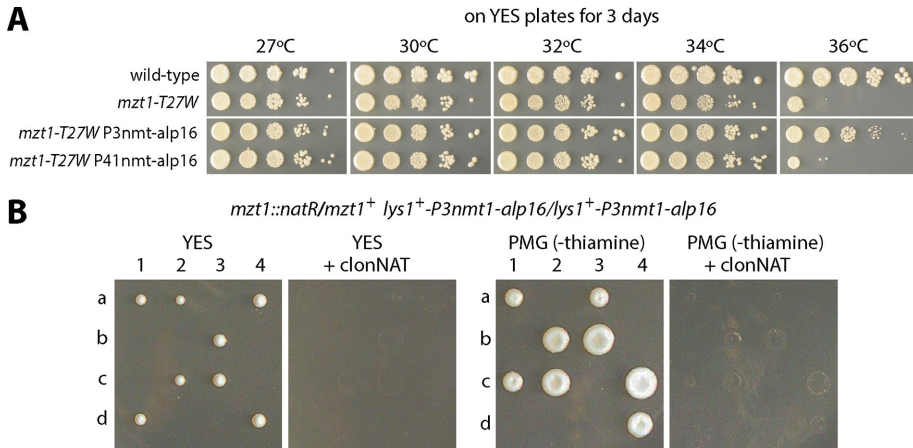


FIGURE 6: $Alp16^{GCP6}$ overproduction suppresses the temperature sensitivity of $mzt1$ mutants but not the lethality of $mzt1$ -null. (A) The temperature sensitivity of $mzt1$ -T27W is rescued by $Alp16^{GCP6}$ overproduction. Serial dilution spot assays of wild-type, $mzt1$ -T27W, $mzt1$ -T27W $lys1^+$ - $P3nmt$ - $alp16$, and $mzt1$ -T27W $lys1^+$ - $P41nmt$ - $alp16$ cells. Cells were spotted onto YES plates and incubated at various temperatures for 3 d. (B) $Mzt1$ is essential for growth in $Alp16^{GCP6}$ -overproducing cells. Diploid cells homologous for $lys1^+$ - $P3nmt$ - $alp16$ and heterozygous for $mzt1$ ($mzt1^+/mzt1\Delta::natR$) were sporulated, and individual spores (a–d) in each ascus (1–4) were dissected on YES to partially repress the expression of $Alp16^{GCP6}$ and on PMG plates to highly induce the expression. Only $mzt1^+$ segregants sensitive to clonNAT grew in both conditions.

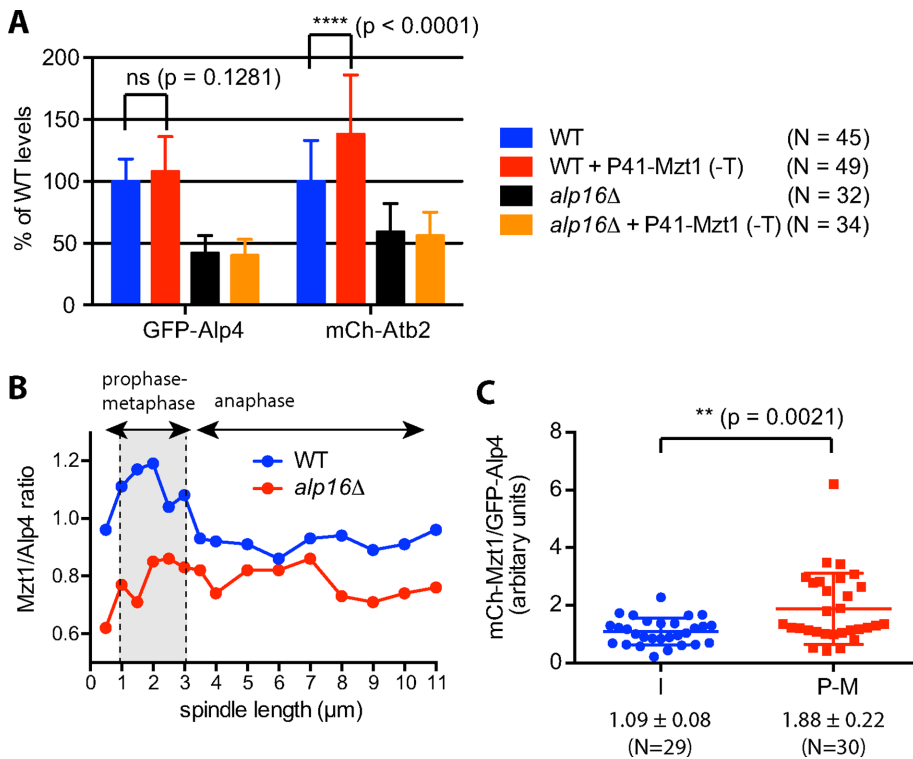


FIGURE 7: $Mzt1$ plays a role in the activation of the γ TuRC ring but not the γ TuSC ring. (A) $Mzt1$ overproduction has no effect on $alp16\Delta$ cells but increases the spindle microtubule level without increasing the level of γ TuC at mitotic SPBs in wild-type cells. $Mzt1$ was overproduced at the $lys1$ locus under the control of $P41nmt$ in the absence of thiamine at 27°C for 22 h in wild-type and $alp16\Delta$ cells containing GFP - $Alp4^{GCP2}$ and mCh - $Atb2$. $Alp4^{GCP2}$ and $Atb2$ levels for spindles <3.5 μ m in length were quantified. (B) The ratio of $Mzt1$ to γ TuC increases at early M phase in wild-type cells. The ratio of $Mzt1$ to $Alp4^{GCP2}$ level at mitotic SPBs was calculated in wild-type and $alp16\Delta$ cells from the data shown in Figure 1, D and E, and plotted against spindle length. (C) The ratio of $Mzt1$ to γ TuC is higher at early M phase than during I phase. The ratio of $Mzt1$ to $Alp4^{GCP2}$ levels at SPBs for spindles <3.5 μ m in length was measured in GFP - $Alp4^{GCP2}$ mCh - $Mzt1$ cells and compared with that at SPBs during interphase.

modification of the γ TuC, which has been suggested by in vitro studies in *S. pombe* (Masuda et al., 1992; Masuda and Shibata, 1996) and shown in vivo in *S. cerevisiae* (Pereira et al., 1998; Vogel et al., 2001; Keck et al., 2011; Lin et al., 2011), or from the mitosis-specific interaction of the γ TuC with regulators required for bipolar spindle assembly, such as kinesins (Prigozhina et al., 2001; Rodriguez et al., 2008; Olmsted et al., 2014) and spindle-anchoring factors (Toya et al., 2007; Yukawa et al., 2015). For instance, kinesin-14 binds to the γ TuRC and establishes spindle bipolarity (Prigozhina et al., 2001; Olmsted et al., 2014). $Alp16$ -specific mitotic phenotypes may be induced through a mechanism involving this interaction.

Alternatively, it may result from differences in the properties of the mitotic and interphase γ TuC receptors. γ TuC receptors can be classified by the N-terminal γ TuSC-binding motifs (Lin et al., 2014). The mitotic receptor $Pcp1$ (Flory et al., 2002; Fong et al., 2010) and the interphase receptor $Mto1$ (Sawin et al., 2004; Lynch et al., 2014) belong to different classes. $Pcp1$ homologues act as a major receptor in *A. nidulans* and human cells, which exhibit $GCP6$ -specific mitotic phenotypes. The γ TuC receptors may define the roles of γ TuRC-specific components in spindle assembly.

Assembly of the γ TuRC ring

We presume that the γ TuRC ring is assembled in the cytoplasm of *S. pombe* cells as shown in other organisms, although there is no direct evidence for its presence. The down-regulation of $GCP4$, $GCP5$, or $GCP6$ disrupts γ TuRC in *Drosophila* and human cells. In contrast, it is not clear for *S. pombe* cells whether the deletion of $Gfh1^{GCP4}$, $Mod21^{GCP5}$, or $Alp16^{GCP6}$ disrupts the γ TuRC ring. Neither gel filtration chromatography (Fujita et al., 2002) nor sucrose gradient centrifugation (Anders et al., 2006; unpublished data) of cell extracts reveals a difference in the distribution pattern of γ TuCs between wild-type cells and $alp16\Delta$ or triple disruptants. These results suggest that γ TuRCs assembled during interphase are unstable or too low in number to be detected by conventional methods using asynchronous cell extracts (Anders et al., 2006). The γ TuRC may become functional at the onset of mitosis when it interacts with the mitotic receptor at the SPBs. We showed that $Alp16^{GCP6}$ expression in the $ghf1\Delta mod21\Delta alp16\Delta$ triple disruptant is sufficient for maintaining γ TuC levels at mitotic SPBs and spindle microtubule levels to an extent similar to wild-type cells (Figure 5). This result raises the

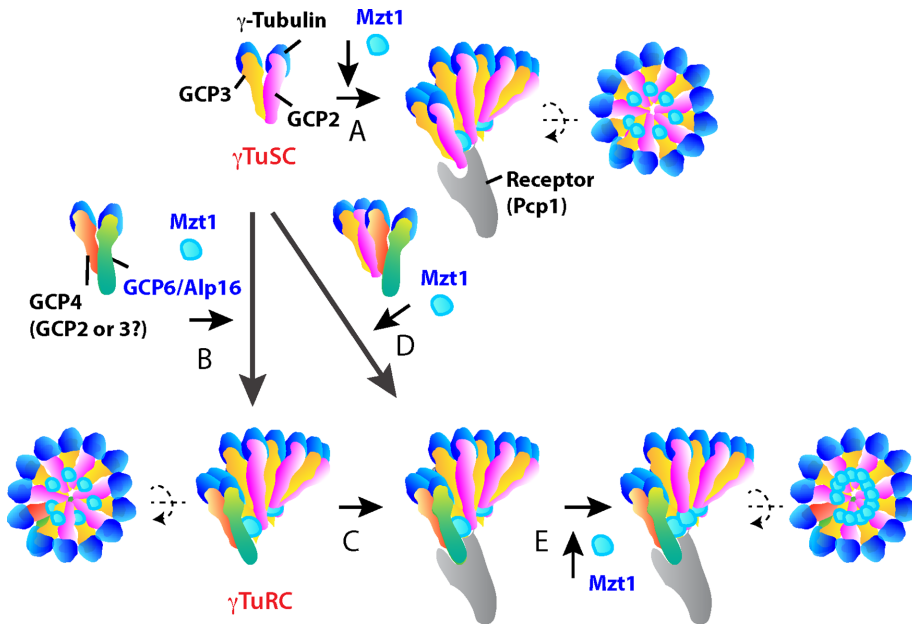


FIGURE 8: Model for the synergistic role of $Alp16^{GCP6}$ and $Mzt1$ in spindle microtubule assembly. A model for the assembly of γ TuSC and γ TuRC rings at mitotic SPBs. (A) The γ TuSC is recruited to Pcp1, the γ TuC receptor at mitotic SPBs, where the γ TuSC ring is assembled. $Mzt1$ is essential for this recruitment. (B) The γ TuRC ring is assembled in the presence of $Alp16^{GCP6}$. $Alp16^{GCP6}$ binds to $Gfh1^{GCP4}$ and to $Alp4^{GCP2}$ or $Alp6^{GCP3}$ in the absence of $Gfh1^{GCP4}$. (C) The γ TuRC ring is recruited to Pcp1 at mitotic SPBs in the presence of $Mzt1$. This recruitment occurs more efficiently than with γ TuSC ring assembly at mitotic SPBs. (D) $Alp16^{GCP6}$ promotes γ TuRC ring assembly at mitotic SPBs in the presence of $Mzt1$. (E) $Mzt1$ fully activates the γ TuRC at mitotic SPBs.

possibility that the γ TuRC is assembled in the cytoplasm without $Gfh1^{GCP4}$ and $Mod21^{GCP5}$ at early M phase (Figure 8B). In the absence of $Gfh1^{GCP4}$ and $Mod21^{GCP5}$, $Alp16^{GCP6}$ may use $Alp4^{GCP2}$ or $Alp3^{GCP3}$ to initiate assembly of the γ TuRC (Figure 8B).

Alternatively, only partial γ TuC rings of various sizes may be assembled in the cytoplasm of *S. pombe* both in the presence and absence of $Alp16^{GCP6}$. The complete γ TuRC ring may be assembled at mitotic SPBs in the presence of $Alp16^{GCP6}$ and $Mzt1$ (Figure 8D). Studies of the *S. cerevisiae* γ TuSC suggest that it is assembled at the SPB but not in the cytoplasm (Kollman *et al.*, 2010; Erlemann *et al.*, 2012; Lin *et al.*, 2014). In *alp16 Δ mod21 Δ gfh1 Δ* cells, only γ TuSCs are present, and therefore the γ TuSC ring presumably is assembled at the MTOCs from γ TuSC or partial γ TuSC rings (Figure 8A). Expression of $Alp16^{GCP6}$ in the triple disruptant would induce $Alp16^{GCP6}$ binding to the γ TuSC (Anders *et al.*, 2006). The $Alp16^{GCP6}$ - γ TuSC complex may promote γ TuRC ring assembly more efficiently at mitotic SPBs when bound to Pcp1 in the presence of $Mzt1$ (Figure 8D). In vitro reconstitution of the γ TuRC ring in the presence of Pcp1 may clarify the role of $Alp16^{GCP6}$ in ring formation.

Synergistic role of $Alp16^{GCP6}$ and $Mzt1$ in spindle microtubule assembly

Although $Alp16^{GCP6}$ promotes recruitment of the γ TuRC to mitotic SPBs and suppresses the temperature sensitivity of the *mzt1* mutant, $Alp16^{GCP6}$ does not act alone as a recruitment factor but in conjugation with $Mzt1$ (Figure 6). The increase in $Alp16^{GCP6}$ expression levels results in an increase in γ TuC levels at mitotic SPBs, which may be due to either $Mzt1$ -dependent attachment of the preformed γ TuRC to mitotic SPBs (Figure 8, B and C), $Mzt1$ -dependent assembly of γ TuRC at mitotic SPBs (Figure 8D), or both. These processes may be more efficient than the γ TuSC ring assembly at

SPBs (Figure 8A). In contrast, elevated $Mzt1$ expression in *alp16 Δ* and wild-type cells does not affect levels of γ TuC at SPBs, suggesting that $Mzt1$ does not act as a rate-limiting factor for γ TuC ring formation at SPBs.

We found that $Mzt1$ overproduction enhances spindle microtubule assembly at mitotic SPBs in wild-type but not in *alp16 Δ* cells. The ratio of $Mzt1$ to $Alp4^{GCP2}$ increases at early M phase in wild-type cells, suggesting that $Mzt1$ has an additional role in the activation of the γ TuRC for spindle microtubule assembly at early M phase (Figure 8E). MOZART1 has been shown to bind GCP3 in *Arabidopsis* and *S. pombe* cells (Janski *et al.*, 2008, 2012; Nakamura *et al.*, 2012; Dhani *et al.*, 2013). $Mzt1$ forms oligomers in vitro, supporting a model in which the $Mzt1$ oligomer stabilizes the γ TuC ring to produce more efficient microtubule nucleation (Dhani *et al.*, 2013). In *Arabidopsis*, MOZART1/GIP1a-containing γ TuC seems to represent a subset of γ TuCs that is more competent for microtubule nucleation (Nakamura *et al.*, 2012), suggesting a role for MOZART1 in γ TuRC activation. We propose that $Mzt1$, presumably in an oligomerized state, fully activates mitotic γ TuRCs for microtubule nucleation. $Alp16^{GCP6}$ and $Mzt1$ are crucial for spatiotemporal regulation of

microtubule nucleation from the γ TuRC and function synergistically at early M phase for efficient bipolar spindle assembly.

MATERIALS AND METHODS

Yeast methods and strains

The *S. pombe* strains used in this study are listed in Supplemental Table S1. Fission yeast media, growth conditions, and manipulations were carried out as previously described (Moreno *et al.*, 1991). For the minichromosome loss assay, strains carrying a minichromosome Ch16 (Niwa *et al.*, 1986) were constructed with the *ade6-M210* background. Because Ch16 had the *ade6-M216* mutation, strains constructed showed the *ade⁺* phenotype due to intragenic complementation, and the loss of Ch16 resulted in the *ade⁻* phenotype detected by red colony or red-sector colony formation on plates containing low concentrations of adenine. Strains carrying Ch16 were grown in yeast extract and supplements (YES) medium overnight, spread on YE agar plates at ~200 cells/plate, and grown at 27°C for 6 d. The ratio (percentage) of the number of red-sector colonies to the number of white- and red-sector colonies combined was calculated as the minichromosome loss rate. For the overproduction of $Alp16^{GCP6}$, $Alp16^{GCP6}$ was expressed under thiamine-repressive promoters at the *lys1* locus in *alp16 Δ* cells. The *alp16⁺* gene was subcloned into pCSU64, 71, and 72 (Chikashige *et al.*, 2004) after GFP was removed from the vectors. pCSU64, 71, and 72 had thiamine-repressive promoters P3nmt, P41nmt, and P81nmt, respectively, for gene expression. The resulting constructs were used for integration of the gene at the *lys1* locus. Cells constructed were incubated at 27°C for 22–24 h in Edinburgh minimal medium 2 (EMM2) supplemented with all growth requirements (amino acids, purines, and pyrimidines) in the presence and absence of 2 μ M thiamine. For the construction of strains carrying proteins

with N-terminal GFP, mCh, and GBP, the promoter region of the gene of interest was PCR amplified and inserted at the *Bam*HI site between a selection marker *kanR* and GFP of pCSS25, mCh of pCSS25-mCh (Masuda *et al.*, 2013), and GBP of pCSS25-GBP. pCSS25-GBP was constructed by replacing the GFP of pCSS25 with GBP. The resulting plasmids were used as templates for PCR-based gene targeting. To examine whether *Alp16^{GCP6}* overproduction rescued the lethality of *mzt1*-null cells, a diploid strain (HR3206) was constructed that was homologous for *lys1⁺-P3nmt-*alp16** and heterozygous for *mzt1* (*mzt1⁺/mzt1Δ::natR*). The diploid cells were sporulated and dissected on YES agar plates to partially suppress the expression of *Alp16^{GCP6}* and on EMM2 and PMG agar plates to fully induce the expression. Pombe medium glutamate (PMG) was the same as EMM2 except that it contained 3.75 g/l L-glutamic acid monosodium salt instead of NH_4Cl . The resistance of viable spores for clonNAT was checked on YES or PMG plates at 100 $\mu\text{g/ml}$.

Fluorescence microscopy

Fluorescence microscope images were obtained using the DeltaVision microscope system (Applied Precision, Seattle, WA) with a CoolSNAP.HQ cooled charge-coupled device camera (Photometrics, Tucson, AZ) equipped with a temperature-controlled chamber (Precision Control, Seattle, WA). Live cells were imaged in a glass-bottom culture dish (MatTek, Ashland, MA) coated with soybean lectin at 27°C. For quantification of fluorescent protein levels, images of 12–14 sections were taken along the z-axis at 0.3- μm intervals. After deconvolution, projection images of maximum intensity were obtained, and maximum fluorescence intensities over the background intensity were used for statistical data analysis. In some experiments, sum intensity images were obtained from five image stacks (a 0.3- μm stack distance) with the stack containing the highest signal intensity in the middle. A 5×5 -pixel (0.5375- μm square) region of interest (ROI) with maximum sum intensity was determined for each mitotic SPB. Sum intensity of two 20×20 -pixel regions around the ROI was used for background subtraction. The data from projection images of maximum intensity were used for all of the analysis, since similar ratios of fluorescence intensity in *alp16Δ* to wild-type cells were obtained using quantification from projection images of the maximum intensity and the sum intensity (Supplemental Figure S2A). Time-lapse imaging was performed by taking images every 1 min for 30 min or every 2 min for 40 min. Spindles formed at 2–10 min after start of observation were selected for analysis. To avoid any effect of photobleaching, cells were observed only under bright field before the fluorescence images were taken. All of the data, except for time-lapse imaging shown in Figures 2 and 3, were obtained from still images (a single time point with z-sectioning).

To test whether an internal fluorescence control is necessary for comparison of GFP levels in wild-type and *alp16Δ* cells, Sid4-mRFP signal intensities at SPBs were measured as internal controls, and GFP intensities relative to mRFP intensities at SPBs (GFP/mRFP ratios) were compared between wild-type and *alp16Δ* cells. The data obtained from the GFP/mRFP ratios were virtually identical to those obtained from GFP intensity in wild-type and *alp16Δ* cells (Supplemental Figure S2, C–I).

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