The negatively charged carboxy-terminal tail of β-tubulin promotes proper chromosome segregation

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ABSTRACT Despite the broadly conserved role of microtubules in chromosome segregation, we have a limited understanding of how molecular features of tubulin proteins contribute to the underlying mechanisms. Here we investigate the negatively charged carboxy-terminal tail domains (CTTs) of α- and β-tubulins, using a series of mutants that alter or ablate CTTs in budding yeast. We find that ablating β-CTT causes elevated rates of chromosome loss and cell cycle delay. Complementary live-cell imaging and electron tomography show that β-CTT is necessary to properly position kinetochores and organize microtubules within the assembling spindle. We identify a minimal region of negatively charged amino acids that is necessary and sufficient for proper chromosome segregation and provide evidence that this function may be conserved across species. Our results provide the first in vivo evidence of a specific role for tubulin CTTs in chromosome segregation. We propose that β-CTT promotes the ordered segregation of chromosomes by stabilizing the spindle and contributing to forces that move chromosomes toward the spindle poles.

INTRODUCTION During mitosis, sister chromatids are separated through a sequence of events orchestrated by a bipolar network of dynamic microtubules known as the mitotic spindle. The spindle assembles from two microtubule nucleation hubs, the spindle poles, which surround the duplicated genome. Microtubules growing out from the spindle poles sample space through cycles of assembly and disassembly until they form linkages that stabilize the spindle and attach to chromatids. The spindle is stabilized by interpolar microtubules (iMTs), a class of microtubules from opposite poles that align in an antiparallel manner, forming extensive lateral contacts. Chromatids attach to kinetochore microtubules (kMTs), a class of microtubules that bind to kinetochores (KTs), multiprotein complexes that assemble at centromeric regions of DNA. These classes of spindle microtubules play unique and important roles that guide chromatid separation.

Sister chromatids must first become bioriented, with the KTs of each sister attaching to kMTs emanating from opposite spindle poles. The progress of biorientation is monitored by signaling pathways that respond to aberrant attachment. Unattached KTs are detected by the spindle assembly checkpoint (SAC), which blocks progression into anaphase (Foley and Kapoor, 2013; Etemad et al., 2015; Tauchman et al., 2015). Syntelic attachments, which arise when both sister chromatids attach to microtubules emanating from the same spindle pole, fail to generate tension and are disrupted by the aurora B/Ipl1 kinase (Lampson and Cheeseman, 2011).

The positions of bioriented sister chromatids are controlled by a balance of forces within the spindle. The strongest evidence for this force balance model comes from studies in budding yeast, which features a relatively simple and well-defined spindle (Winey and Bloom, 2012). Spindle microtubules generate outwardly directed forces that pull each sister toward opposite spindle poles. Major contributors to outward forces include the microtubule motors kinesin-14 and kinesin-5, which organize and generate forces on iMTs, respectively, to push the spindle poles apart (Saunders and Hoyt, 1992; Saunders et al., 1997; Hepperla et al., 2014). Outward forces are opposed by inwardly directed forces that hold sister chromatids...
together. Major contributors to inward forces include loops of pericentric chromatin that are bound together by cohesin (Bouck and Bloom, 2007; Stephens et al., 2013; Chacon et al., 2014). This balance of forces sets the size of the spindle and prepares sister chromatids to separate and move to the spindle poles at anaphase.

How are spindle microtubules regulated to support these various roles? The best-studied role for spindle microtubules is in KT attachment. KTs attach to microtubules through multiple binding modules. The Ndc80 complex binds microtubules through two modules—calponin homology domains in the Ndc80 and Nuf2 subunits, and an unstructured, positively charged N-terminal tail domain in Ndc80 (Wigge and Kilmartin, 2001; Wei et al., 2007; Ciferri et al., 2008; Alushin et al., 2010; Sundin et al., 2011). The Ndc80 tail enhances binding through electrostatic interaction with the microtubule surface and is critical for regulating attachment. When sisters attach to microtubules from the same spindle pole, the microtubule–KT interface is disrupted by the aurora B/Ipl1 and Mps1 (Cheeseman et al., 2009; Alushin et al., 2012; Sarangapani et al., 2013; Zaytsev et al., 2015). In budding yeast, the role of the Ndc80 tail appears to overlap with the Dam1 complex, which interacts with and recruits the Ndc80 complex to microtubule plus ends (Lampert et al., 2010). Like Ndc80, Dam1 binds to microtubules through electrostatic interactions, and it is targeted for phosphorylation by aurora B/Ipl1 and Mps1 (Cheeseman et al., 2002; Westermann et al., 2005; Shimogawa et al., 2006). Mutants impairing either Ndc80’s tail or Dam1 partially disrupt chromosome segregation, but combining these mutations produces severe defects (Demirel et al., 2012; Lampert et al., 2013; Kalantzaki et al., 2015). The complexity of the microtubule–KT interactions underscores the importance of the regulatory mechanisms that promote proper attachment.

In contrast to the KTs, a mechanistic understanding of how microtubules contribute to this highly regulated process is lacking. Tubulin proteins exhibit molecular diversity in genetically encoded amino acid sequences, as well as posttranslational modifications. Nearly all of this diversity is found in the negatively charged carboxy-terminal tail (CTT) regions of α- and β-tubulin, which decorate the surface of microtubules (Roll-Mecak, 2015). CTTs are known to support electrostatic interactions with KTs. For example, biochemically removing CTTs disrupts binding to Ndc80 and Dam1 in vitro (Westermann et al., 2005; Ciferri et al., 2008). However, the roles of CTTs in a complex cellular environment are undefined. In this study, we use mutants that alter or ablate CTTs of α- and β-tubulin in budding yeast to test the role of CTTs in vivo and gain new insight into chromosome segregation mechanisms. Our results demonstrate a specific role for the CTT of β-tubulin in promoting the fidelity of chromosome segregation and timely cell cycle progression. Mutant cells lacking β-CTT exhibit defects in KT positioning and disorganized spindle microtubules. We identify a short region of negatively charged amino acid residues in β-CTT that are necessary and sufficient for high-fidelity chromosome segregation and provide evidence that the role of β-CTT is conserved across species.

RESULTS

β-CTT promotes proper chromosome segregation

We first tested whether CTTs of α- or β-tubulin are important for chromosome segregation by measuring the loss of a nonessential chromosome fragment (Figure 1A; Spencer et al., 1990). Mutants lacking β-CTT (tub2-430Δ) exhibit a fivefold increase in chromosome loss events compared with wild type (WT; Figure 1B). In contrast, mutants either lacking the major α-CTT (tub1-442Δ) or all α-CTTs (tub1-442Δ tub3-442Δ) are indistinguishable from WT in this assay. Mutants lacking both α-CTT and β-CTT exhibit a rate of chromosome loss that is similar to mutants lacking β-CTT alone (Figure 1B). Therefore β-CTT specifically promotes chromosome segregation.

To examine whether β-CTT might function in a common pathway with KT protein complexes, we used the chromosome loss assay to test for genetic interactions. Yeast mutants that disrupt the Ndc80 tail perturb chromosome segregation and exhibit additive effects when combined with Dam1 mutants (Kemmler et al., 2009; Demirel et al., 2012; Kalantzaki et al., 2015). We found that double mutants combining Ndc80 tail truncation (n0c80-112Δ) with β-CTT...
truncation lost chromosomes at a rate similar to β-CTT single mutants but greater than ndc80-112Δ single mutants (Figure 1B). In contrast, double mutants combining Dam1 impairment (dam1-1, at a semipermissive temperature of 25°C) with β-CTT truncation lose chromosomes at a significantly higher rate than either single mutant (Figure 1B). This additive effect is reminiscent of the genetic interaction between ndc80-112Δ and dam1-1 and indicates that cells depend on the function of both β-CTT and the N-terminal tail of Ndc80 when Dam1 is impaired.

β-CTT is necessary for timely progression through mitosis

If chromosome loss in mutants lacking β-CTT arises from defects in spindle assembly, then these mutants might exhibit a SAC-dependent delay in cell cycle progression. We performed a series of experiments to test this prediction. First, we used liquid growth assays to show that mutants lacking β-CTT exhibit a 20% increase in doubling time compared with WT controls and mutants lacking all α-CTTs (tub1-442Δ, tub3-442Δ; Figure 2A). Second, we examined individual synchronized cells, measuring the time from bud emergence to separation in cells released from START. Cells expressing Pds1-13myc were collected at 15-min intervals, prepared for Western blots, and probed with myc antibodies. (D) Pds1-13myc signal at each time point normalized to t = 0. Values are averages from three experiments. Error bars are SEM.

In contrast, β-CTT mutants exhibit variable Nufl2-GFP localization, including single clusters, two WT-like clusters, and two clusters positioned very close to the SPBs (Figure 3B and Supplemental Figure S1B). dam1-765 mutants consistently exhibit two clusters of Nufl2-GFP very close to the SPBs, as expected (Figure 3C and Supplemental Figure S1C). This initial result suggests that KT position may be more variable in β-CTT mutants.

We used several approaches to quantify differences in Nufl2-GFP localization across populations of preanaphase cells. First, we measured the volume within the cell that is occupied by Nufl2-GFP, based on single-time point confocal Z-series imaging. This analysis shows that Nufl2-GFP samples a larger volume within β-CTT mutant cells than in WT controls (Figure 3D). Of importance, total Nufl2-GFP signal per cell is similar in both WT and mutant strains; therefore changes in Nufl2-GFP localization in β-CTT mutants are attributable to changes in position rather than changes in protein levels or number of KTs (Figure 3E). In separate experiments, we found that the localization of an inner KT component, the centromeric histone variant Cse4/CENP-A, is similarly altered in mutants lacking β-CTT (Supplemental Figure S1, D and E).

We next examined Nufl2-GFP localization across different spindle lengths. KTs only resolve into two clusters once the spindle reaches a sufficient length (Marco et al., 2013). In the course of our experiments, we observed that shorter spindles are more abundant in β-CTT mutants (Figure 3F), raising the possibility that differences in Nufl2-GFP localization could be caused by differences in spindle length. To test this possibility, we compared the percentages of cells exhibiting a bilobed distribution of Nufl2-GFP at different spindle lengths. This analysis shows that a similar percentage of WT and β-CTT mutant cells exhibit bilobed Nufl2-GFP, regardless of spindle length (Figure 3G). In contrast, more dam1-765 mutant cells exhibit bilobed Nufl2-GFP across all spindle lengths examined (Figure 3G). In addition to comparing the separation of KT clusters, we also examined the positions of KT clusters along the spindle axis. In WT spindles shorter than 1100 nm, Nufl2-GFP localizes to a single lobe at the center of the spindle (Figure 3H). In WT spindles longer than 1100 nm, Nufl2-GFP signal shifts away from the spindle center and toward the poles, consistent with a bilobed distribution.
β-CTT promotes chromosome segregation

Nuf2-GFP signal gradually resolves into a bilobed distribution until the two lobes rapidly move apart, indicating the onset of anaphase (Figure 3K). In preanaphase β-CTT mutant cells, Nuf2-GFP resolves into a bilobed distribution, but this state is interrupted by prolonged periods of collapse into a single, unresolved lobe (Figure 3K, arrow). At anaphase, the rate of Nuf2-GFP separation in β-CTT mutants is slower than in WT cells; this is consistent with our previous finding that the rate of spindle elongation is slower in β-CTT mutants (Aiken et al., 2014). These results support our conclusion that β-CTT is necessary to maintain KT position in the spindle.

β-CTT is necessary to align KTs with the spindle axis

To identify the cause of KT positioning defects, we imaged a single pair of sister KTs by labeling chromosome IV with GFP inserted 2 kb from the centromere (CENIV). Imaging CENIV sisters clearly reveals the bioriented state—GFP-labeled CENIV resolves into two foci only when attached to kMTs from opposite spindle poles (Figure 4, A and B, and Supplemental Figure S2, A and B; Pearson et al., 2004). Surprisingly, we found that a greater percentage of Nuf2-GFP signal gradually resolves into a bilobed distribution until the two lobes rapidly move apart, indicating the onset of anaphase (Figure 3K). In preanaphase β-CTT mutant cells, Nuf2-GFP resolves into a bilobed distribution, but this state is interrupted by prolonged periods of collapse into a single, unresolved lobe (Figure 3K, arrow). At anaphase, the rate of Nuf2-GFP separation in β-CTT mutants is slower than in WT cells; this is consistent with our previous finding that the rate of spindle elongation is slower in β-CTT mutants (Aiken et al., 2014). These results support our conclusion that β-CTT is necessary to maintain KT position in the spindle.
Preanaphase \( \beta \)-CTT mutant cells exhibit bioriented CENIV compared with preanaphase WT cells (Figure 4C). It is important to note that our method did not resolve CENIV foci separated by <470 nm; therefore the “unseparated” population may include sisters that are bioriented but closely apposed. Nevertheless, \( \beta \)-CTT is clearly not necessary for biorientation.

We next used our CENIV imaging assay to compare the positions of bioriented sister centromeres in the three-dimensional (3D) space of the spindle. To determine whether centromeres are displaced further from the SPB-SPB axis, we calculated the alignment angles of each CENIV sister relative to the SPB-SPB axis (see Materials and Methods). These angles are significantly increased in \( \beta \)-CTT mutants compared with WT controls (70 ± 2° [mean ± SEM] in \( \text{tub2-430A} \) vs. 58 ± 3° for WT; Figure 4D and Supplemental Figure S2C). kMT lengths, that is, the distances between each CENIV and the nearest SPB, are similar in \( \beta \)-CTT mutants and WT controls, as are the distances between CENIV sisters (Figure 4E and Supplemental Figure S2D, D and E). Spindle lengths are slightly shorter in \( \beta \)-CTT mutants (\( \rho = 0.06 \); Figures 3F and 4E and Supplemental Figure S2F). These data suggest that \( \beta \)-CTT may normally act to confine bioriented sister centromeres near the SPB-SPB axis.

The positions of bioriented sister centromeres are determined by a balance of outward forces that pull them apart (tensile attachments to kMTs and forces from iMTs on the SPBs) and inward forces hold them together (cohesin, condensin; Bouck and Bloom, 2007; Gardner et al., 2008; Stephens et al., 2013; Chacon et al., 2014). To determine how \( \beta \)-CTT contributes to forces in the spindle, we combined the \( \beta \)-CTT truncation mutation with cohesin mutations that relax inward forces. Disrupting pericentric cohesin with an \( \text{mcm21} \) null mutation increases the percentage of preanaphase cells with bioriented CENIV, increases the distance between sister centromeres, and increases spindle length (Figure 4, C and E; Stephens et al., 2011). We reasoned that if centromere-positioning defects in \( \beta \)-CTT mutants are primarily caused by weaker outward forces, then simultaneously relaxing inward forces with \( \text{mcm21} \) might rescue these defects and lead to intermediate phenotypes. We found that although double mutants combining \( \text{tub2-430A} \) with \( \text{mcm21} \) do rescue CENIV alignment angles, spindle lengths are increased similarly to those in \( \text{mcm21} \) single mutants (Figure 4, D and E). In addition, double mutants exhibit significant increases in kMT length and distance between CENIV sisters and a greater percentage of cells with separated CENIV than either single mutant alone (Figure 4, C and E). Therefore disrupting cohesin appears to exacerbate some spindle defects in \( \beta \)-CTT mutants.

\( \beta \)-CTT regulates microtubule organization during spindle assembly

To examine how changes in microtubules might accompany changes in KT position, we mapped the organization and lengths of spindle microtubules using electron tomography. We first compared microtubules in preanaphase spindles, which we identified based on spindle length. Preanaphase WT cells exhibit ~32 short microtubules (~600 nm) and ~8 longer microtubules (>800 nm; Figure 5, A and B). These correspond to kMTs that attach to each KT and iMTs.
that interdigitate to form antiparallel overlaps that stabilize the bipolar spindle, respectively (Winey et al., 1995). Preanaphase spindles in β-CTT mutants differ from WT controls in several respects. First, the distance between SPBs is decreased (Supplemental Figure S3). This is consistent with our light microscopy results and our previous findings from time-lapse imaging (Figure 3F; Aiken et al., 2014).

Second, the two spindle halves are often misaligned and exhibit few interdigitating microtubules (Figure 5C and Supplementary Figure S3). This is reminiscent of an early stage of spindle assembly, before the bipolar state is stabilized (Winey and O'Toole, 2001). Three of four tomograms of preanaphase β-CTT mutant spindles exhibit a clear misalignment of spindle halves, and therefore it is unlikely that this represents a transient intermediate. Instead, β-CTT mutants may linger in this state.

Third, β-CTT mutant spindles contain more microtubules and exhibit differences in microtubule lengths. In general, β-CTT mutant spindles contain a greater frequency of shorter (<200 nm) microtubules and fewer longer microtubules (Figure 5E). Many of these microtubules deviate far from the SPB-SPB axis (Figure 5C and Supplementary Figure S3). The changes in microtubule length and alignment in β-CTT mutants may explain changes in KT position that we observed by light microscopy, including aberrant Nu1-2-GFP distribution and deviation of KTs from the SPB-SPB axis (Figures 3 and 4).

Fourth, β-CTT mutant spindles contain microtubule “fragments” that are not connected to SPBs and exhibit flared morphology at both ends (Figure 5C and Supplementary Figure S3, colored yellow). Fragments vary in length and number per cell, as do their location in the spindle. We did not find microtubule fragments in WT cells, and, to our knowledge, they have not been previously reported.

Although β-CTT mutants exhibit microtubule defects in preanaphase spindles, we did not observe defects in anaphase spindles. Anaphase cells for both WT and the β-CTT mutant exhibit ∼16 very short (<100 nm) microtubules from each SPB, along with a small number of much longer microtubules that overlap in the center of the spindle (Figure 5, F–I, and Supplemental Figure S3). These represent kMTs and iMTs, respectively. The lengths and number of these microtubules are not significantly different between mutant and WT. We conclude that β-CTT is important for regulating microtubule organization during spindle assembly.

Mapping the region of β-CTT that is important for chromosome segregation

Finally, we sought to gain insight into the mechanism of β-CTT function by mapping the amino acids that are important for its role in chromosome segregation. Yeast β-tubulin is enriched for negatively charged amino acids at residues 431–438 (Figure 6A). We term this region the “acidic patch.” Restoring these eight amino acids rescues the defects of tub2-430Δ, including chromosome segregation, growth rate, and viability in the absence of SAC activity (Figure 6, B and C; unpublished data). Restoring the acidic patch and the next seven residues (tub2-445Δ) also rescues the defects of tub2-430Δ (Figure 6, B and C). These results indicate
that the acidic patch is sufficient for β-CTT function in chromosome segregation.

Although the sequence of the acidic patch is not strictly conserved across β-tubulin isotypes, the enrichment of negatively charged residues is a common feature (Figure 6D and Supplemental Table S1). To determine whether negative charge is necessary for function, we replaced glutamate and aspartate residues in β-tubulin with glutamines. Neutralizing these residues in β-tub2-CTT region of β-tubulin (Supplemental Table S1). (E) Genetic interactions with SAC mutants. Tenfold dilution series of strains indicated at left were spotted to rich medium or rich medium supplemented with 10 μg/ml benomyl and grown at 30°C.

DISCUSSION

Our study provides the first in vivo evidence of tubulin’s contribution to the electrostatic interactions that regulate the mitotic spindle. We demonstrate that negatively charged amino acid residues within the CTT region of β-tubulin promote proper chromosome segregation. Ablating these residues or neutralizing the charges of their side chains increases chromosome segregation errors and makes cells reliant on the SAC for viability. We propose that the principal role of β-CTT is to organize spindle microtubules to stabilize the spindle and promote the ordered separation of sister chromatids.

The role of β-CTT in spindle assembly is supported by our findings, as well as previous studies. Time-course experiments with synchronized cells show that mutants lacking β-CTT delay the formation of bipolar spindles (Aiken et al., 2014). In the present study, we find that preanaphase spindles are shorter in β-CTT mutants, based on distributions of spindle lengths in asynchronous populations of cells (Figure 3F) and specifically in metaphase cells with bioriented CENIV distributions (Figure 4E). These new results, combined with our previous findings, indicate that β-CTT may contribute to both the early stages of spindle assembly and the subsequent maintenance of spindle stability in metaphase. Our tomographic analyses provide insight into the specific role of β-CTT in these contexts. In contrast to wild-type controls, β-CTT mutants show misalignment of the two halves of the preanaphase spindle and a concomitant lack of interdigitating iMTs (Figure 5 and Supplemental Figure S3). We analyzed four β-CTT
mutant cells with preanaphase spindles (determined by the distance between SPBs and the lengths of kMTs), and all four exhibit these defects. These characteristics are reminiscent of WT cells during the early stages of spindle assembly and suggest that β-CTT may be necessary for either efficient transit through early stages of spindle assembly or the stabilization of the metaphase spindle (Winiey and O’Toole, 2001). Moreover, the kMTs in the mutant cells appear to be grossly disorganized (Figure 5C). The disorganization of kMTs is consistent with the disorganization we observed at the levels of kinetochores and sister centromeres (Figures 3 and 4). Of importance, structural defects in β-CTT mutants are specific to preanaphase spindles; we did not observe similar defects in tomograms of anaphase spindles (Figure 5 and Supplemental Figure S3). Determining the molecular mechanism of β-CTT function during spindle assembly will require further study; however, we speculate that this mechanism may involve electrostatic interactions with positively charged microtubule-binding proteins that promote the alignment and/or cross-linking of interdigitating iMTs.

How could microtubule disorganization lead to defective KT positioning and, ultimately, chromosome missegregation? Our results suggest that β-CTT is not necessary to biorient individual sister KTs but is necessary to cluster KTs near the SPBs and maintain the alignment of KTs along the SPB-SPB axis (Figures 3 and 4). Of importance, the defects we observe in β-CTT mutants are quite different from those observed in mutants that alter KT-microtubule binding (e.g., dam1-1765; Figure 3I; Shimogawa et al., 2006) or inhibit biorientation (e.g., ipil1 or stu2 mutants; Marco et al., 2013). We propose that β-CTT may contribute to both the generation of outward forces on KTs and the maintenance of kMTs along the spindle axis (Figure 7). Our analysis of tub2-430Δ mcm21Δ double mutants supports this model. If the spindle defects in β-CTT mutants were attributable to weakened outward forces alone, then simultaneously weakening inward forces with the cohesin mutant would be expected to restore the balance of forces in the spindle, and double mutants should therefore exhibit intermediate phenotypes relative to either single mutant alone. This relationship was demonstrated for mutants that disrupt kinesin-5 motors (Bouck and Bloom, 2007). We do find that tub2-430Δ mcm21Δ double mutants rescue KT alignment along the SPB-SPB axis, indicating that imbalanced spindle forces contribute to KT positioning defects (Figure 4D). However, tub2-430Δ mcm21Δ double mutants also exhibit longer spindles that are similar to those in mcm21Δ single mutants (Figure 4E) and an even greater percentage of cells with separated CENIV than for either single mutant (Figure 4C). Therefore loss of the cohesin network (mcm21Δ) is synergistic with loss of β-CTT mutants, indicating that β-CTT also has an important role in clustering centromeres along the spindle axis. Taken together, these results show that whereas outward forces are affected in β-CTT mutants, the mechanism could also involve defective kinetochore microtubule organization. Another question that arises is whether the distortions of spindle structure in β-CTT mutants alter normal functions like the surveillance of KT attachment by the SAC. This could reconcile the paradox of how β-CTT mutants experience cell cycle delay and chromosome segregation errors despite the apparent efficiency of biorientation.

An alternative but not mutually exclusive model is that β-CTT could regulate electrostatic interactions at the KT-microtubule interface. We find no evidence of deficient biorientation in β-CTT mutants, indicating that interactions with tubulin CTTs are not necessary for KTs to attach to microtubules in budding yeast. This result highlights molecular differences between the yeast and mammalian systems, in which electrostatic interactions play an important role in regulating KT attachment (DeLuca et al., 2006; Guimaraes et al., 2008). One possible explanation for these differences is that yeast possess other unique modes for binding KTs to microtubules, such as the Dam1 complex, which might act in parallel to electrostatic interactions with tubulin CTTs. Consistent with this, our epistasis experiments in Figure 1B reveal strongly additive chromosome segregation defects in double mutants that simultaneously ablate β-CTT and impair the Dam1 complex. Therefore fully functional Dam1 appears to become necessary for chromosome segregation when β-CTT is missing, consistent with the two acting in parallel. Previous studies of yeast mutants that lack the N-terminal tail of Ndc80 found a similar relationship with the Dam1 complex (Demirel et al., 2012). Collectively, these results support a model in which the Ndc80 tail and β-CTT act together to support a KT-microtubule interface that acts in parallel to the Dam1 complex.

Finally, our findings raise the question of whether molecular changes at the level of β-CTT, either genetically encoded or post-translational, may provide a mechanism for regulating mitotic spindle function. CTT sequences are highly divergent across species and across tubulin isoforms within a species (Supplemental Table S1). Furthermore, CTTs are major targets of posttranslational modifications that dramatically alter CTT structure and charge (Roll-Mecak, 2015). Together these differences create a highly variable molecular landscape at the microtubule surface. Our results indicate that sequence differences between β-tubulin isoforms do not alter spindle function as long as the CTT region contains a sufficient quorum of negatively charged amino acid residues (Figure 6). Whether posttranslational modifications of β-CTT alter spindle function remains an open and important question.

**MATERIALS AND METHODS**

Chemicals and reagents were from Fisher Scientific (Pittsburgh, PA) and Sigma-Aldrich (St. Louis, MO), unless stated otherwise.

**Yeast strains and manipulation**

General yeast manipulation, media, and transformation were performed by standard methods (Amberg et al., 2005). A detailed list of strains is provided in Supplemental Table S2. Mutant alleles of...
TUB1, TUB2, and TUB3 were generated at the native chromosomal loci as described in Aiken et al. (2014), ndc80-112Δ, which lacks the N-terminal tail encoded by the first 112 amino acids, was generated by removing codons 2–112 from the native NDC80 locus. dam-1-1 was generated by creating a C111Y substitution mutation at the native DAM1 locus, recreating the previously identified dam1-1 allele (Jones et al., 1999). dam-1-765 was generated by cloning the dam-1-765 allele (a gift from Trisha Davis, University of Washington, Seattle, WA), containing the S221F substitution, into a plasmid containing the Schizosaccharomyces pombe his5 marker, creating plasmid pJM0283. The dam1-765::SpHis5 cassette was then amplified by PCR and integrated into the native locus, replacing the native DAM1 allele. Strains expressing CENIV labeled with GFP were derived from those described in Brito et al. (2010). All other fluorescent fusions were integrated at the endogenous chromosomal loci.

**Chromosome loss assay**

Retention of a nonessential chromosome fragment carrying an ochre-suppressing allele of the SUP11 tRNA, which suppresses premature termination of ade2-101 translation, was measured as described in Koshland and Hieter (1987). Cultures were grown overnight at 30°C in synthetic –URA dropout medium. The saturated cultures were diluted 100-fold, and cells were counted using a hemocytometer. Two hundred cells were plated on yeast extract/peptone/dextrose (YPD) and grown for 3–4 d at 30°C before being moved to 4°C for 2 d for color development. The plates were scanned at 600-dpi resolution in 24-bit color. Colonies were counted either by hand or by using a custom-made MATLAB program. The half-red-sectorized colonies were divided by the total number of colonies to determine chromosome loss frequency. Each strain was tested in at least three separate experiments, with at least 1000 colonies scored per experiment.

**Liquid growth assay**

Cells were grown in 3 ml of YPD overnight at 30°C and diluted 50-fold into fresh medium. The diluted cultures were then aliquoted into a 96-well plate, with eight replicates per experiment, and incubated at 30°C with single orbital shaking in a CytoLine3 plate reader (BioTek, Winooski, VT). The OD$_{590}$ was measured every 5 min for 24 h. Doubling time was calculated by fitting the growth curves to a nonlinear exponential growth curve (GraphPad Prism version 6.0; GraphPad, San Diego, CA).

**Budding duration analysis**

Budding duration was determined by time-lapse microscopy of α-factor–synchronized cultures grown to mid log phase (WT, 2 h; tub2-430Δ, 3 h). Cultures were released from arrest by washing twice and resuspending cells in fresh medium supplemented with protease, then mounted on slides and imaged every 15 min for 135 min. Images were collected on a Nikon Ti-E microscope (Nikon, Melville, NY) equipped with a 100×/numerical aperture (NA) 1.49 CFI 60 Apochromat objective, PEKA light engine (Lumencore, Beaverton, OR), and scientific complementary metal-oxide semiconductor camera (ORCA-Flash 4.0 LT; Hamamatsu Photonics, Shizuoka, Japan) using NIS Elements software (Nikon).

**Time course of Pds1 levels**

Cells were grown to early log phase at 30°C in 100 ml of YPD and arrested with two pulses of α-factor for 90 and 60 min, respectively. The cells were pelleted, washed with water, resuspended into 130 ml of fresh YPD, and returned to the 30°C shaking incubator. We collected 15-ml culture samples at 15-min intervals, which were washed, pelleted, and lysed by bead beating. The total protein concentration of the clarified lysate was determined by Bradford assay, and all samples were normalized. Samples were run on SDS-PAGE, transferred to nitrocellulose membrane, and blocked overnight at 4°C. Membranes were probed with mouse α-myc (MA1-980; 1:1000; Thermo Scientific, Waltham, MA) and goat α-mouse-800 (926-32210; 1:10,000; LI-COR Biosciences, Superior, NE) and imaged on an Odyssey Imager (LI-COR Biosciences). Intensities of Pds1-myc bands were quantified using ImageJ (National Institutes of Health, Bethesda, MD). Pixel intensities were measured across an equivalent area for each Western blot lane within a time course, and the intensities of these regions were divided by the intensity at t = 0 to obtain a ratio of each time point to the initial level.

**Microscopy and image analysis**

Images were collected on a Nikon Ti-E microscope equipped with a 100×/1.45 NA CFI Plan Apo objective, piezoelectric stage (Physik Instrumente, Auburn, MA), spinning-disc confocal scanner unit (CUSt; Yokogawa, Tokyo, Japan), 488- and 561-nm lasers (Agilent Technologies, Santa Clara, CA), and an electron-multiplying charge-coupled device (CCD) camera (iXon Ultra 897; Andor Technology, Belfast, United Kingdom) using NIS Elements software. Cells were grown asynchronously to early log phase in nonfluorescent medium, mounted on a slab of 2% agarose, and sealed beneath a coverslip with Vaseline/lanolin/paraffin at 1:1:1. During acquisition, the temperature of the stage was 25°C. The Z-series consisted of 21 images separated by 300 nm.

Preanaphase cells were identified in asynchronous populations based on spindle length, which was determined by measuring the distance between SPBs labeled with Spc110-RFP. The distribution of all spindle lengths from asynchronous cells shows a peak that reflects metaphase spindle length, and the mean values for WT (1.8 μm) and tub2-430Δ (1.6 μm) cells were consistent with the average spindle length just before anaphase onset, which we previously defined using time-lapse imaging (Figure 3F; Aiken et al., 2014). We therefore defined “preanaphase” as bipolar spindles shorter than the mean of the total population.

The volume occupied by Nu2-GFP was determined from single-time point confocal stacks of cells expressing Nu2-GFP and Spc110-DSRed from the native loci. Preanaphase cells were identified and segmented from the field using a custom ImageJ macro. After segmentation, a low-bandpass Butterfield filter and semi-automated Otsu intensity thresholding application were used to identify Nu2-GFP signal and generate a binary mask. Masks were then converted to regions of interest (ROIs) using connected-components pixel labeling. The total number of pixels in the ROI was then converted into micrometers cubed using an empirically derived pixel calibration in x and y (57 nm/pixel) and a z-depth of the stack (300 nm). We validated this approach by comparing the sum of Nu2-GFP pixel values measured in each cell and found that the data exhibited a normal distribution that was not different between WT and ΔACTT mutant cells.

The distribution of Nu2-GFP along the spindle axis was measured by using a custom MATLAB program to determine the centroid coordinates of the spindle poles (labeled with Spc110-DSRed). Briefly, image stacks were compressed to a sum intensity projection and then processed using a Laplacian filter and Otsu intensity thresholding to generate binary masks. Two-dimensional ROIs were defined using connected-components pixel labeling, and the x,y-coordinates of the centroids of these ROIs were calculated using MATLAB. Spindle lengths were calculated between centroids in MATLAB using a distance formula. Each sum intensity projection
image was rotated using calculated angle that aligned the spindle along a horizontal axis. Images were then cropped to 10 pixels (570 nm) on either side of the SPBs and 7 pixels above and below the SPB-SPB axis (800 nm total). Line scans of NuF2-GFP intensity were then generated by summing pixel values in each 14-pixel column along the SPB-SPB axis.

Centroids of CENIV-GFP and Spc110-RFP were determined by first segmenting individual cells and then using a custom MATLAB program to determine the centroid coordinates. Briefly, image stacks were processed using Laplacian filter and Otsu intensity thresholding to generate binary masks. Three-dimensional ROIs were defined in z-stacks using connected-components pixel labeling, and the x,y,z-coordinates of the centers of these ROIs were calculated using MATLAB. kMT lengths were calculated in MATLAB using a 3D distance formula, based on the distances from the centroid of a bioriented sister CENIV and the nearest SPB. kMT angles were calculated using the inverse cosine of kMT length divided by the distance from CENIV to the SPB-SPB axis.

Electron tomography
Cells were prepared for electron microscopy as described in Giddings et al. (2001). Briefly, aliquots from log-phase liquid cultures were collected onto a 0.45-μm Millipore filter (EMD Millipore, Billerica, MA) by vacuum filtration. The cells were then frozen using a Wohlfend Compact 02 high-pressure freezer (Wohlfend, Sennwald, Switzerland), freeze substituted in 0.25% glutaraldehyde and 0.1% uranyl acetate in acetone, and embedded in Lowicryl HM20 resin. Serial sections (250 nm) were collected onto Formvar-coated slot grids and poststained with 2% uranyl acetate followed by lead citrate. Colloidal gold particles were affixed to the sections to serve as fiducial markers for alignment.

Tomography was performed essentially as described in O’Toole et al. (2002). Dual-axis tilt series were collected using a Tecnai F20 or F30 microscope (FEI, Hillsboro, OR) using the SerialEM program for data acquisition (Mastronarde, 2005). Tilt-series data were acquired from serial sections at a pixel size of 1.2 nm using a Gatan (Pleasanton, CA) CCD camera. Tomographic volumes were computed using the IMOD software package (Kremer et al., 1996; Mastronarde, 1997). Volumes from three to or four serial sections were joined, and the spindle microtubules were modeled using the 3 dmod program of the IMOD software package. Microtubule lengths were computed from the model contour data.

Benomyl sensitivity
Cells were grown in YPD to saturation at 30°C, and a 10-fold dilution series of each was spotted to either YPD or YPD supplemented with 10 μg/ml benomyl. Plates were grown at 30°C for 2 d.

Sequence logo
A probability-weighted Kullback–Leibler sequence logo of β-CTT residues 426–445 was created using the Seq2Logo server (www.cbs.dtu.dk/biotools/Seq2Logo; Thomsen and Nielsen, 2012) based on the distance from CENIV to the SPB-SPB axis.

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REFERENCES
Aiken J, Sept D, Costanzo M, Boone C, Cooper JA, Moore JK (2014).
Genome-wide analysis reveals novel and discrete functions for tubulin carboxy-terminal tails. Curr Biol 24, 1295–1303.
Alushin GM, Munsinpally V, Matson D, Tooley J, Stukenberg PT, Nogales E (2012). Multimodal microtubule binding by the Ndc80 kinetochore complex. Nat Struct Mol Biol 19, 1161–1167.
Alushin GM, Ramey VH, Pasqualato S, Ball DA, Grigorieff N, Musacchio A, Nogales E (2010). The Ndc80 kinetochore complex forms oligomeric arrays along microtubules. Nature 467, 805–810.
Amberg DC, Burke D, Strathern J (2005). Methods in Yeast Genetics, Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press.
Bouch DC, Bloom K (2007). Pericentric chromatin is an elastic component of the mitotic spindle. Curr Biol 17, 741–748.
Brito IL, Yu HG, Amon A (2010). Condensins promote coorientation of sister chromatids during meiosis I in budding yeast. Genetics 185, 55–64.
Chacon JM, Mukherjee S, Schuster BM, Clarke DJ, Gardner MK (2014). Pericentromere tension is self-regulated by spindle structure in metaphase. J Cell Biol 205, 313–324.
Cheeseman IM, Anderson S, Jwa M, Green EM, Kang J, Yates JR, Chan CS, Drubin DG, Barnes G (2002). Phospho-regulation of kinetochore-microtubule attachments by the Aurora kinase Ipl1p. Cell 111, 163–172.
Ciferri C, Pasqualato S, Srecpani E, Varette G, Santaguida S, Dos Reis G, Maiolica A, Polka J, De Luca GJ, De Wulf P, et al. (2008). Implications for kinetochore-microtubule attachment from the structure of an engineered Ndc80 complex. Cell 133, 427–439.
Cohen-Fix O, Peters JM, Kirschner MW, Koshland D (1996). Anaphase initiation in Saccharomyces cerevisiae is controlled by the APC-dependent degradation of the anaphase inhibitor Pds1p. Genes Dev 10, 3081–3093.
DeLuca GJ, Gall WE, Ciferri C, Cimini D, Musacchio A, Salmon ED (2006). Kinetochore-microtubule dynamics and attachment stability are regulated by Hecl. Cell 127, 969–982.
Demirel PB, Keyes BE, Chatterjee M, Remington CE, Burke DJ (2012). A redundant function for the N-terminal tail of Ndc80 in kinetochore-microtubule interaction in Saccharomyces cerevisiae. Genetics 192, 753–756.
Ettamad B, Kuijt TE, Kops GJ (2015). Kinetochore-microtubule attachment is progression by kinesin-5 motor-mediated disassembly of longer kinetochore microtubules. Cell 133, 894–906.
Giddings THJ, O’Toole ET, Morpew M, Mastronarde DC, McIntosh JR, Winey M (2001). Using rapid freeze and freeze-substitution for the preparation of yeast cells for electron microscopy and three-dimensional analysis. Methods Cell Biol 67, 27–42.
Goshima G, Yanagida M (2000). Establishing biorientation occurs with precocious separation of the sister kinetochores, but not the arms, in the early spindle of budding yeast. Cell 100, 619–633.
Guimaraes GJ, Dong Y, McEwen BF, Deluca JG (2008). Kinetochore-microtubule attachment relies on the disordered N-terminal tail domain of Hecl. Curr Biol 18, 1778–1784.
He X, Asthana S, Sorger PK (2000). Transient sister chromatid separation and elastic deformation of chromosomes during mitosis in budding yeast. Cell 101, 763–775.

Hepperla AJ, Willey PT, Coombs CE, Schuster BM, Gerami-Nejad M, McClellan M, Mukherjee S, Fox J, Winey M, Odde DJ, et al. (2014). Minus-end-directed kinesin-14 motors align antiparallel microtubules to control metaphase spindle length. Dev Cell 31, 61–72.

Jones MH, Bachant JB, Castillo AR, Giddings TH, Winey M (1999). Yeast Dam1p is required to maintain spindle integrity during mitosis and interacts with the Mps1p kinase. Mol Biol Cell 10, 2377–2391.

Kalantzaki M, Kitamura E, Zhang T, Mino A, Novak B, Tanaka TU (2015). Kinetochore-microtubule error correction is driven by differentially regulated interaction modes. Nat Cell Biol 17, 421–433.

Kemmler S, Stach M, Knapp M, Ortiz J, Pfannstiel J, Ruppert T, Lechner J (2009). Mimicking Ndc80 phosphorylation triggers spindle assembly checkpoint signalling. EMBO J 28, 1099–1110.

Koshland D, Hieter P (1987). Visual assay for chromosome ploidy. Methods Enzymol 155, 351–372.

Kremer JR, Mastronarde DN, McIntosh JR (1996). Computer visualization of three-dimensional image data using IMOD. J Struct Biol 116, 71–76.

Lampert F, Hornung P, Westermann S (2010). The Dam1 complex confers microtubule plus end-tracking activity to the Ndc80 kinetochore complex. J Cell Biol 189, 641–649.

Lampert F, Mieck C, Alushin GM, Nogales E, Westermann S (2013). Molecular requirements for the formation of a kinetochore-microtubule interface by Dam1 and Ndc80 complexes. J Cell Biol 200, 21–30.

Lampson MA, Cheeseman IM (2011). Sensing centromere tension: Aurora B and the regulation of kinetochore function. Trends Cell Biol 21, 133–140.

Lampson MA, Cheeseman IM (2011). Sensing centromere tension: Aurora B and the regulation of kinetochore function. Trends Cell Biol 21, 133–140.

Lampson MA, Cheeseman IM (2011). Sensing centromere tension: Aurora B and the regulation of kinetochore function. Trends Cell Biol 21, 133–140.

Lampson MA, Cheeseman IM (2011). Sensing centromere tension: Aurora B and the regulation of kinetochore function. Trends Cell Biol 21, 133–140.

Lampson MA, Cheeseman IM (2011). Sensing centromere tension: Aurora B and the regulation of kinetochore function. Trends Cell Biol 21, 133–140.