INTRODUCTION

Microtubules are involved in essential cellular processes including chromosome movement during mitosis, vesicular trafficking, cellular polarization, and mRNA targeting. Microtubules are dynamic structures, i.e., at polymer mass steady state, individual microtubules are observed growing, depolymerizing, or remaining in a paused or attenuated state, in which no changes in microtubule length are observed (Mitchison and Kirschner, 1984). The dynamics of microtubules are controlled by a number of factors including microtubule-associated proteins (Drechsel et al., 1992; Kowalski and Williams, 1993; Dhamodharan and Wadsworth, 1995; Hamill et al., 1998), catastrophe-promoting proteins (reviewed in Walczak, 2000), motor proteins (reviewed in Hunter and Wordeman, 2000), and the GTPase activity of tubulin (reviewed in Erickson and O’Brien, 1992).

The subunit of microtubules, tubulin, is a heterodimer consisting of α- and β-tubulin. A number of biochemical studies have suggested that tubulin cysteine residues play roles in tubulin assembly into microtubules. For example, chemical modification of cysteine residues results in inhibition of tubulin assembly in vitro (Ludueña and Roach, 1991) and GTP hydrolysis by tubulin (Mejillano et al., 1996). Cysteine residues can also be covalently cross-linked to the exchangeable site guanine nucleotide (Shivanna et al., 1993; Jayaram and Haley, 1994; Bai et al., 1999), to a colchicine analogue (Bai et al., 1996) and to other antimitotic agents (Bai et al., 1989; Shan et al., 1999). Previously, we used site-directed mutagenesis to investigate the roles of the six cysteine residues in Saccharomyces cerevisiae Tub2p (β-tubulin) in the structure and function of tubulin (Gupta et al., 2001). Of particular interest were the tub2-C354S and tub2-C354A mutations that produced phenotypes characteristic of greatly increased microtubule stability, including cold stability of microtubules and increased benomyl resistance (Gupta et al., 2001). The increased stability could reflect alterations in the intrinsic properties of the microtubule or altered binding of regulatory proteins. The ability to correlate in vivo and in vitro dynamics allows us to distinguish these possibilities.

The budding yeast S. cerevisiae contains few microtubules compared with most other eukaryotic cells (Huffaker et al.,...
allowing one to dissect the contribution of microtubules and their dynamic properties to specific cellular events. The cytoplasmic microtubules are required for positioning the nucleus to the neck before anaphase onset, and contribute to spindle elongation in anaphase. The nuclear microtubules form the bipolar spindle and are required for chromosome segregation. It has been proposed that the highly dynamic nature of microtubules contributes to a search and capture mechanism of cytoplasmic microtubules at the site of bud growth during G1 (Carminati and Stearns, 1997; Shaw et al., 1997b) and of nuclear microtubules at the kinetochores before chromosome segregation (Holy and Leibler, 1994; Winiey et al., 1995; O’Toole et al., 1999; Maddox et al., 2000). However, it has not been possible to distinguish the role of microtubule dynamics from the essential role of microtubules for cellular viability.

In this report we utilize two tub2-C354 mutant strains that exhibit altered microtubule dynamics in vivo and in vitro. We find that cytoplasmic microtubule dynamics are greatly reduced. In addition, the number of cytoplasmic microtubules is reduced to approximately one per spindle pole body. Although this stable microtubule is able to locate the incipient bud site, the spindle is not aligned correctly along the mother-bud axis. Moreover, tubulin turnover in the nuclear spindle microtubules is reduced in the mutant strains. The defect in microtubule dynamics restricts the extent and rate of spindle elongation during anaphase. The cell cycle regulation of microtubule dynamics is also altered. In wild-type cells, microtubules are most dynamic in G1, whereas in the tub2-C354 mutants, G1 dynamics in unbudded cells are reduced relative to budded cells. These studies reveal important roles of microtubule regulatory proteins throughout the cell cycle.

MATERIALS AND METHODS

Yeast Strains

The yeast strains used in this study were MG1 (MATa, leu2Δ1, trplΔ63, his3-917, URA3/ura3-52, tub2-His2), which contains a His2 tag at the C terminus of Tub2p, and the MG1 mutant strain MG1-C354S (MATa, leu2Δ1, trplΔ63, his3-917, URA3/ura3-52, tub2-His2-C354S; Gupta et al., 2001). Also, created for this study were the strains FY1-GFP (MATa, LEU2::GFP-TUB1::leu2Δ1, trplΔ63, his3-917, URA3/ura3-52, TUB1, TUB2), which is FY1 (Davis et al., 1993) containing the GFP-Tub1p fusion protein (described below) and the FY1-GFP mutant strains: FY1-GFP-C354A (MATa, LEU2::GFP-TUB1::leu2Δ1, trplΔ63, his3-917, URA3/ura3-52, TUB1, tub2-C354A) and FY1-GFP-C354S (MATa, LEU2::GFP-TUB1::leu2Δ1, trplΔ63, his3-917, URA3/ura3-52, TUB1, tub2-C354S), which are FY1-GFP containing the respective tub2p-C354 mutations (described below).

Construction of the GFP-Tub1p Strains

The plasmid pMG3 encoding GFP-Tub1p under control of the TUB1 promoter was cloned by transforming the GFP-TUB1 construct from pAFS125 (A. Straight, Harvard University, Cambridge, MA) into the multiple cloning site of the yeast integrating plasmid pRS405. The 3100-base pair kpnI-Sacl fragment from pAFS125 containing GFP-TUB1 and the endogenous TUB1 promoter was combined with the 924-base pair kpnI-KasI and 447-base pair KasI-Sacl fragments from pRS405 in a three-way ligation reaction. When digested at the single KasI site, pMG3 integrates at the LEU2 locus.

The strain FY1-GFP was created by transforming FY1 with linearized pMG3. The mutant strains FY1-GFP-C354A and FY1-GFP-C354S were constructed by transforming FY1-GFP with the transplacement fragment from pCS3 (Sage et al., 1995a) containing a tub2-C354A or tub2-C354S mutation, respectively. All DNA manipulations, isolation of transformants, and verification of mutations were done as described previously (Gupta et al., 2001).

In Vivo Microtubule Dynamics: Microscopic Imaging

Cells were grown to midlog phase and mounted as described previously (Yeh et al., 1995; Maddox et al., 2000) using SD-complete media (0.67% yeast nitrogen base without amino acids, 2% glucose, 0.5% casamino acids), and 50 µg/ml each uracil, tryptophan, and adenine. The equipment and techniques for imaging GFP-fusion proteins have been described in detail (Shaw et al., 1997a). Cells were grown and maintained at 23°C during image acquisition. The typical acquisition protocol acquired five z-series fluorescent images at 0.75-µm axial steps and a single differential interference contrast (DIC) image corresponding to the central fluorescent image. Time-lapse image series, which were typically acquired at 8-s intervals for wild-type microtubules, were usually acquired every 20 s for tub2-C354 mutant microtubules in order to extend the duration of the analysis. For microtubule dynamics analysis, time-lapse series lasted between 6 and 60 min. For spindle dynamics and cell cycle analysis, image sets were typically acquired every 60 s and the total series could cover >2 h before significant photobleaching occurred.

In Vivo Microtubule Dynamics: Image Analysis and Quantification

Yeast cells were segregated into four categories for in vivo microtubule dynamics analysis. Unbudded cells were single cells with no bud growth visible by DIC microscopy. Cells were classified as small-budded from the time an emerging bud became visible by DIC microscopy until duplicated spindle pole body separation became visible by fluorescence microscopy. The separation of duplicated spindle pole bodies was evident by two distinct dots of GFP-Tub1p fluorescence replacing the previously singular dot. Budded cells that had a bipolar spindle <2 µm in length were classified as preanaphase cells. Budded cells were identified as anaphase cells from the time the mitotic spindle had elongated to >2 µm in length until spindle midzone separation became visible.

Only microtubules and spindles whose entire length lay within the series of z-focus plane images were used for analysis. If possible, multiple cells from the same time-lapse series or multiple microtubule images from a single cell were analyzed. In the case of the tub2-C354A mutant, and to a lesser extent the tub2-C354S mutant, cells were seen with deviations in cell morphology (Gupta et al., 2001). Cells with individual characteristics not representative of the strain as a whole were not used for microtubule dynamics analysis.

At each time point, microtubules were analyzed by identifying the tip and the base of the microtubule in their respective z-plane images. The Cartesian coordinates and the z-plane separation distance of the microtubule ends were used to calculate the three-dimensional length of the microtubule. Each series of time-lapse images was analyzed three separate times as independent sets of measurements, which were used to construct microtubule lifetime history plots using the averages of the three length measurements.

Microtubule dynamics rates were calculated by linear regression analysis of the lifetime history plots. Growth and shortening events are defined as a set of at least four consecutive time-points with an R² value ≥0.85 and a length excursion ≥ 0.6 µm. Pause events are defined as at least four consecutive time-points during which length change was < ±0.2 µm. Because at least four consecutive time-points were considered in linear regression analysis, all growth, shortening, and attenuation events in wild-type and tub2-C354 mutant cells lasted longer than 24 and 60 s, respectively. Catastrophe is defined as a transition to shortening after a period of growth or pause. Similarly, rescue is defined as a transition to growth following a period of pause or depolymerization. Brief periods in some of
the time-lapse series remained unclassified because they did not meet the criteria set for time duration, length of excursion, or statistical significance. The proportion of time spent in each phase was calculated by dividing the sum of the time spent in each phase by the total evaluated time for all microtubules observed. The frequencies of catastrophe and rescue were calculated by dividing the number of events observed by the total evaluated time. SDs were calculated as the frequencies of catastrophe or rescue divided by the square root of the number of events observed (Walker et al., 1988). Microtubule dynamics was calculated as the total number of tubulin dimers exchanged per microtubule end (using 1690 dimers/μm microtubule) considering the proportion of time spent in growth, shortening, or pause phase (Toso et al., 1993).

The number of cytoplasmic microtubules was determined by GFP-Tub1p fluorescence using single time point z-series images. Fluorescent intensity line scan analysis (MetaMorph software) was performed on a three-pixel line along the length of microtubules that were contained within a single z-focal plane image. The gray-scale intensity values of each line scan were scaled to demonstrate relative changes in intensity.

**Spindle Elongation, Position, and Orientation**

Spindle length during elongation, from the outside edges of the spindle poles, was calculated in the same manner described for in vivo microtubule analysis and quantification. Spindle position was determined by measuring the distance from the outside edge of the bud-proximal spindle pole body to the center of the bud neck in two-dimensional z-series composite images ~1 min before the spindle elongated to >2 μm. To determine the angle of orientation for proanaphase spindles, time-lapse sequences were selected which displayed the complete spindle (~2 μm long) for at least 10 min. At 1-min intervals, the angle of a line intersecting both spindle poles was determined relative to the mother-daughter axis of the cell in two-dimensional z-series composite images. A total of 10, 13, and 15 spindle images were used to determine spindle orientation for wild-type, tub2-C354S, and tub2-C354A mutant strains, respectively.

**FRAP Experiments**

Fluorescence recovery after photobleaching (FRAP) experiments and statistical analysis were carried out as described in Maddox et al. (2000). Five 350-ms exposure images were acquired at each time point through focus z-section with 30-s-2 min intervals. The mitotic half-spindle was photobleached using a 35-ms laser exposure. The ratio of bleached/unbleached region after recovery R, first-order rate constant k, and half-time to recovery (t1/2) were calculated at the following average time points: 2 min (FY41-GFP), 17.6 min (FY41-GFP-C354S), and 18.6 min (FY41-GFP-C354A).

**Purification of Yeast Tubulin**

Yeast strains MGY1 and MGY1-C354S were grown in 34 L of YPD medium (1% yeast extract, 2% peptone, 2% glucose) in 2–20 L carboys. After overnight growth at 30°C with vigorous aeration, the medium was supplemented with an additional 2% glucose, and the pH was adjusted to ~6.5 with 4 M NaH4OH. Growth was continued for an additional 5 h, and cells harvested in a Sharples Type T-1P continuous flow centrifuge (Pennwalt Corp., Warminster, PA). Approximately 500 g of packed, wet weight cells were obtained.

Wild-type cells were suspended in 100 ml of H-PEM (100 mM Pipes, 2 mM EGTA, 10 mM MgSO4, pH 6.9) and stored at 4°C overnight. (In our previous article we found that it takes at least 6 h to completely depolymerize yeast cellular microtubules at 4°C [Gupta et al., 2001].) Because incubation of MGY1-C354S cells at 4°C induces the formation of cold-stable tubulin polymers (Gupta et al., 2001), these cells were suspended in H-PEM and processed immediately for tubulin purification. Immediately before purification, the total volume of H-PEM was brought to 750 ml and made 5 mM in dithiothreitol, 2 mM in phenylmethylsulfonyl fluoride, and 0.1 mM in GTP. The cells were lysed by passing the suspension 10 times through a Microfluidizer 110-Y (Microfluidics Corp., Newton, MA) using an ice-cooling coil. This method produced >95% cell breakage.

The initial stages of the purification procedure were based on the procedures of Barnes et al. (1992) and Davis et al., (1993). All steps were done at 4°C. The lysed cells were centrifuged at 10,000 × g for 10 min, and the resulting supernatant was centrifuged at 100,000 × g for 60 min. The 100,000 × g supernatant was made 10% in glycerol and 0.16 M in NaCl. A 400-ml bed volume of DE52 (Whatman Inc., Fairfield, NJ), equilibrated with H-PEM + 0.16 M NaCl, was added as a 70% slurry, and the suspension was stirred gently for 45 min.

The DE52 was washed twice by centrifugation (7000 × g for 10 min) and resuspension in 600 ml FMG (100 mM Pipes, 1 mM MgSO4, 10% glycerol, pH 6.9) containing 0.16 M NaCl, after which it was suspended in 300 ml of the same solution and loaded into a 30 × 5-cm column. After gravity packing, the column was washed with an additional 75 ml of the same solution. The majority of proteins were not retained by the resin. Remaining proteins were eluted with PMG25 (FMG with 25 mM Pipes) containing 0.5 M NaCl. Fractions containing tubulin, determined by FRAP assay (Bradford, 1976), were pooled and brought to 20 mM imidazole using a 500 mM solution, and the pH was adjusted to 7.4 with NaOH. A 5-ml bed volume of Ni-NTA resin (Qiagen, Valencia, CA), washed twice in 50 ml PMG25 containing 20 mM imidazole and 0.5 M NaCl, pH 7.4, was added to the pooled fractions as a 50% slurry. The suspension was stirred gently for 45 min, and the Ni-NTA resin was collected by centrifugation for 7 min at 3000 × g. The resin was washed twice with 25 ml of the same solution followed by 15 ml of a solution containing 25 mM Pipes, 1 mM MgSO4, 20 mM imidazole, pH 7.4. The resin was finally suspended in 5 ml of the latter solution and loaded into a 20 × 0.5-cm column. Proteins were eluted with 350 mM imidazole, pH 6.9, using a flow rate of 0.2 ml/min, and 0.5-ml fractions were collected. Tubulin eluted immediately following one void volume. Fractions containing >0.75 mg/ml tubulin were pooled and dialyzed twice for 45 min against 300 volumes of 10 mM Pipes, 0.1 mM EGTA, 0.1 mM MgSO4, 50 μM GTP, pH 6.9, in a 10,000 MWCO Slide-A-Lyzer Cassette (Pierce Chemical Co., Rockford, IL). Fractions of 0.2–0.75 mg/ml tubulin were pooled and dialyzed against 0.1 M NaCl, 0.1 M MgSO4, 20 mM EGTA, pH 6.9, then passed through a 1000-cm2 membrane (Millipore Corp., Bedford, MA) at 13°C to achieve steady state. Images of microtubules were recorded for no longer than 90 min after steady state was achieved. Thirty-two wild-type and 16 tub2p-C354S microtubules were analyzed using the RTM software kindly provided by E. D. Salmon (University of North Carolina). Length measurements were recorded every 30–45 s over the lifetime of the microtubule or, in the case of rapid depolymerization events, with the time of event exceeding 3–4 s. For all growth phases the average growth rate >1.8 μm/h, and shortening rates were >3.6 mm/h. For tub2p-C354S microtubules, all growth phases had time-averaged growth rates >1.8 μm/h, and pauses were >10 min with time-averaged
length changes of $<\pm 0.6$ $\mu$m/h. For tub2p-C354S microtubules, shortening rates were $>180$ $\mu$m/h. Microtubule dynamic parameters were defined and calculated as described for in vivo analysis. Polymer mass steady state was verified by assembling 1.4 mM tubulin under identical conditions with the same concentration of purified axonemes. At time points between 15 and 120 min, aliquots were analyzed by a microtubule sedimentation assay. The amount of polymer was constant through 120 min of incubation for both wild-type and tub2p-C354S tubulin.

**RESULTS**

A Single Stable Cytoplasmic Microtubule Facilitates Spindle Positioning

Wild-type cells typically contain several cytoplasmic microtubules as visualized with GFP-Tub1p. Unbudded and small-budded wild-type cells contained an average of $2.7 \pm 0.9$ and $3.2 \pm 1.2$ cytoplasmic microtubules, respectively. The majority of cells transformed with tub2-C354S or tub2-C354A in place of wild-type TUB2 contained only a single cytoplasmic microtubule emanating from the spindle pole body (Figure 1A). Tub2-C354S mutants contained $1.1 \pm 0.3$ and $1.1 \pm 0.4$, whereas tub2-C354A mutants contained $1.3 \pm 0.5$ and $1.1 \pm 0.4$ cytoplasmic microtubules in unbudded and small-budded cells, respectively (p < 0.001 for all vs. wild-type). During anaphase, $1.0 \pm 0.3$ microtubules emanated from each of the duplicated spindle pole bodies in both tub2-C354 mutants (p < 0.003 for both mutants vs. $1.4 \pm 0.3$ for wild-type). The number of cytoplasmic microtubules in wild-type and tub2-C354 mutants was statistically indistinguishable during preanaphase (Figure 1A). We used line scan analysis to determine whether the structure in the mutants was indeed one microtubule or perhaps a bundle of a few microtubules (Figure 1, B–I). The constancy of the intensity throughout the structure is indicative of a single microtubule. The decrease in intensity at the tip is observed in wild-type and mutant microtubules.

Cytoplasmic microtubules in un budded cells pivot around the spindle pole body, and this movement may facilitate localization of the incipient bud site. The single microtubule in tub2-C354 mutant cells, which underwent infrequent and minor length changes (discussed later), exhibited a similar type of pivoting from the spindle pole body as wild-type microtubules (Figure 2B). By the time an
emerging bud was visible by transmitted light microscopy, the microtubule was stationary, with the tip extending into the bud. As the bud grew, the microtubule remained oriented in the bud (Figure 3). The microtubule shortened, with the microtubule tip remaining in the bud, thereby positioning the spindle adjacent to the bud neck (Figure 3).

To determine the fidelity of spindle positioning, we measured the distance from the bud-proximal spindle pole body to the center of the bud neck ~1 min before anaphase B (spindle length > 2 μm). In wild-type cells, the spindle was located 0.83 ± 0.60 μm (n = 9) from the bud neck. Spindles were located 1.27 ± 0.70 μm (n = 13) and 0.93 ± 0.76 μm (n = 15) from the bud neck in tub2-C354S and tub2-C354A mutant cells, respectively. Despite the larger average spindle-to-neck distance in the mutant cells, the differences between the strains were statistically indistinguishable (p = 0.13 and 0.74 for wild-type vs. tub2-C354S and tub2-C354A, respectively).

Although the spindle was positioned near the bud neck at the time of spindle elongation, the mutant cells displayed defects in spindle orientation before anaphase onset. In wild-type cells, the preanaphase spindle was aligned within 30° of the mother-daughter axis 87% of the time. The preanaphase spindle was greater than 30° offset from the mother-daughter axis 51% of the time in tub2-C354S cells and 28% of the time in tub2-C354A cells.

**Cytoplasmic Microtubule Dynamics in tub2-C354 Mutant Cells**

All parameters of microtubule dynamics were severely depressed in tub2-C354 mutant GI cells (Table 1). Dynamic rates of tub2-C354 mutant microtubules were statistically different from the corresponding rates for wild-type microtubules (p < 0.001). Growth and shrinkage rates were decreased by 83-92% compared with wild-type microtubules. The tub2-C354 mutant microtubules had transition frequencies that were 2-5% of those for wild-type microtubules and spent the great majority of the time in the paused state...
Table 1. In vivo cytoplasmic microtubule dynamic rates and frequencies*

<table>
<thead>
<tr>
<th></th>
<th>Wild-type</th>
<th>tub2-C354$S$</th>
<th>tub2-C354$A$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Growth rate (μm/min)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>1.66 ± 0.54 (31)</td>
<td>0.29 ± 0.16 (4)</td>
<td>0.13 ± 0.02 (4)</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>1.33 ± 0.52 (11)</td>
<td>0.29 ± 0.13 (5)</td>
<td>NA$^b$ (0)</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>1.30 ± 0.42 (28)</td>
<td>0.23 ± 0.11 (9)</td>
<td>NA$^b$ (0)</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>1.31 ± 0.41 (15)</td>
<td>0.40 ± 0.20 (24)</td>
<td>0.12 ± 0.04 (8)</td>
</tr>
<tr>
<td>Shrinkage rate (μm/min)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>2.66 ± 0.77 (34)</td>
<td>0.46 ± 0.21 (6)</td>
<td>0.28 ± 0.03 (3)</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>2.24 ± 0.54 (9)</td>
<td>0.31 ± 0.23 (9)</td>
<td>0.27 ± 0.18 (6)</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>1.98 ± 0.68 (30)</td>
<td>0.42 ± 0.18 (9)</td>
<td>0.15 (1)</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>1.89 ± 0.74 (18)</td>
<td>0.49 ± 0.21 (15)</td>
<td>0.31 ± 0.20 (2)</td>
</tr>
<tr>
<td>Catastrophe frequency (min$^{-1}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>0.52 ± 0.10 (28)</td>
<td>0.012 ± 0.005 (5)</td>
<td>0.012 ± 0.007 (3)</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>0.45 ± 0.15 (9)</td>
<td>0.036 ± 0.014 (7)</td>
<td>0.036 ± 0.016 (5)</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>0.46 ± 0.10 (24)</td>
<td>0.024 ± 0.009 (7)</td>
<td>0.006 (1)</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>0.25 ± 0.08 (10)</td>
<td>0.072 ± 0.021 (12)</td>
<td>0.012 ± 0.008 (2)</td>
</tr>
<tr>
<td>Rescue frequency (min$^{-1}$)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>0.26 ± 0.09 (14)</td>
<td>0.006 ± 0.004 (3)</td>
<td>0.012 ± 0.007 (3)</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>0.15 ± 0.08 (3)</td>
<td>0.018 ± 0.009 (4)</td>
<td>&lt;0.006 (0)$^b$</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>0.21 ± 0.07 (11)</td>
<td>0.036 ± 0.012 (9)</td>
<td>&lt;0.005 (0)$^b$</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>0.22 ± 0.07 (9)</td>
<td>0.108 ± 0.025 (18)</td>
<td>0.024 ± 0.011 (5)</td>
</tr>
<tr>
<td>Dynamicity (dimers/s)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>49.1</td>
<td>0.99</td>
<td>0.52</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>35.9</td>
<td>3.48</td>
<td>1.96</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>33.1</td>
<td>1.49</td>
<td>0.25</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>25.4</td>
<td>5.90</td>
<td>1.26</td>
</tr>
</tbody>
</table>

* In wild-type cells 10 to 25 microtubules were analyzed. In tub2-C354$S$ cells, 8 to 16 microtubules were analyzed. In tub2-C354$A$ cells 5 to 15 microtubules were analyzed. Results are reported as the mean ± 1 SD. The number of events is in parentheses.

No events were observed during the total time analyzed.

(Table 1 and 2). The dynamicity of the tub2-C354$S$ microtubules was 2% that of wild-type microtubules, and the tub2-C354$A$ mutation produced microtubules with 1% the dynamicity of wild-type microtubules (Table 1). In 700 min of observation none of the microtubule depolymerizations in the mutant unbudded cells resulted in the microtubule becoming <1 μm in length. By contrast, in 53 min of observation 22 of the 25 wild-type microtubules analyzed in unbudded cells underwent length excursions < 1 μm.

The parameters of microtubule dynamics in the mutant cells were altered during the cell cycle in a different manner than in wild-type cells (Table 1). For example, the growth and shrinkage rates in wild-type cells were lower in anaphase than in unbudded cells, but in the mutants these parameters did not change significantly during the cell cycle. On the other hand, microtubules in the mutants showed much more dramatic changes in transition frequencies, proportion of time spent in each phase, and dynamicity. For example, in the tub2-C354$S$ mutant there was a 6-fold difference in catastrophe and an 18-fold difference in rescue frequency between unbudded and anaphase cells. In addition, the pattern of cell cycle–dependent microtubule dynamics modulation differed in the mutant cells. Wild-type microtubules were most dynamic in unbudded cells and became progressively less dynamic throughout the cell cycle, decreasing in dynamicity by 49% between unbudded and anaphase cells. In the mutant cells, microtubules were least dynamic in unbudded and preanaphase cells, and dynamicity was stimulated as much as fourfold and sixfold in small-budded and anaphase cells, respectively. In both mutants, the increased microtubule dynamicity in small-budded cells was effected by a 3-fold increase in catastrophe frequency and a 10-fold increase in the proportion of time spent in catastrophe.

Table 2. Time distribution of in vivo microtubule dynamic phases

<table>
<thead>
<tr>
<th>Time spent in growth (%)</th>
<th>Wild-type</th>
<th>tub2-C354$S$</th>
<th>tub2-C354$A$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unbudded cells</td>
<td>47.5</td>
<td>3.3</td>
<td>7.5</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>50.7</td>
<td>5.6</td>
<td>0.0$^b$</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>40.5</td>
<td>9.8</td>
<td>0.0$^b$</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>29.3</td>
<td>31.5</td>
<td>25.8</td>
</tr>
<tr>
<td>Time spent in shrinkage (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>35.2</td>
<td>3.5</td>
<td>2.2</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>25.8</td>
<td>33.1</td>
<td>23.2</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>31.4</td>
<td>7.1</td>
<td>2.2</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>26.1</td>
<td>17.0</td>
<td>2.8</td>
</tr>
<tr>
<td>Time spent attenuated (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unbudded cells</td>
<td>17.3</td>
<td>93.2</td>
<td>90.3</td>
</tr>
<tr>
<td>Small-budded cells</td>
<td>23.5</td>
<td>61.4</td>
<td>76.8</td>
</tr>
<tr>
<td>Preanaphase cells</td>
<td>28.3</td>
<td>83.0</td>
<td>97.8</td>
</tr>
<tr>
<td>Anaphase cells</td>
<td>44.6</td>
<td>51.4</td>
<td>71.4</td>
</tr>
</tbody>
</table>

* The data come from the experiments described in Table 1.

$^b$ No growth events were observed.
spent shrinking, which resulted in overall shortening of the cytoplasmic microtubule. Conversely, rescue frequency was increased to about twice that of the catastrophe frequency in anaphase cells, and in the case of the tub2-C354S mutant, was accompanied by a 10-fold increase in the percent time spent growing, resulting in net elongation of the cytoplasmic microtubule during anaphase.

Representative lifetime histories of microtubules in wild-type and mutant cells are presented in Figure 4. The difference between wild-type and mutant microtubules is readily apparent. Although microtubules in the wild-type strain underwent several transitions within 300 s, those in the mutant strains were mostly paused for up to 3500 s. Additionally, the data in Figure 4 demonstrate that the mutant

Figure 4. Cytoplasmic microtubule dynamic behavior in wild-type and tub2-C354 mutant cells. Representative lifetime history plots constructed from three-dimensional length measurements (see MATERIALS AND METHODS) are presented. The length axis is identical for all plots. Note the scale of the time axis is different for wild-type and mutant microtubules. (A) Microtubules in unbudded wild-type cells undergo several length excursions in <300 s. (B) Microtubules in unbudded tub2-C354 mutant cells are relatively nondynamic for as long as 3500 s. (C) In small-budded tub2-C354 mutants, dynamicity is increased nearly fourfold relative to unbudded cells and microtubules undergo net depolymerization. (D) In preanaphase cells, tub2-C354 mutant microtubules are again relatively nondynamic. (E) During anaphase, dynamicity is up to sixfold higher than in unbudded cells and tub2-C354 mutant microtubules undergo net polymerization. Panels B–E show representative lifetime history plots of either tub2-C354A or tub2-C354S microtubules.
microtubules typically underwent net disassembly in small-budded cells, were relatively nondynamic in unbudded and preanaphase cells, and underwent net growth in anaphase cells. Tub2-C354 mutant cells completed anaphase with a single cytoplasmic microtubule on each spindle pole body. These microtubules had elongated during anaphase and remained long and stable during the subsequent G1 phase.

### Spindle Microtubule Dynamics

The mitotic spindle, consisting of kinetochore and interpolar microtubules, is typically disassembled within a few minutes of the completion of anaphase (Maddox et al., 2000; Pearson et al., 2001). The kinetochore microtubules depolymerize during anaphase A (Winery et al., 1998), and the interpolar microtubules separate at the midzone and depolymerize toward the spindle poles during telophase (Maddox et al., 2000). In the tub2-C354 mutants, the mitotic spindle failed to depolymerize after spindle separation in the midzone, and a remnant half-spindle was visible as a fluorescent tuft extending from the spindle pole body (Figure 2). Therefore, the mutant unbudded and small-budded cells contained remnant spindle microtubules from the previous mitosis. The spindle microtubules in the mutants typically depolymerized before spindle assembly in the ensuing cell cycle.

To determine if the tub2-C354 mutations affect spindle microtubule dynamics, we measured the spindle tubulin turnover rate in these mutants. Measurement of the dynamics of individual microtubules in the yeast spindle is not possible because of the high density of microtubules within the mitotic spindle. However, FRAP of GFP-Tub1p–labeled spindle microtubules can be used to measure the turnover of tubulin within the mitotic spindle. Previous studies have shown that spindle microtubules in wild-type cells turnover with a half-life of approximately 1 min (Maddox et al., 2000). This turnover is likely due to the dynamic instability of kinetochore microtubules growing and shortening at their plus ends.

To measure the turnover of GFP-Tub1p in the mutant spindle, we photobleached half of the mitotic spindle and then monitored the recovery of the bleached half-spindle fluorescence intensity and the decrease of the unbleached half-spindle fluorescence intensity (Figure 5). The tub2-C354S mutant showed a first-order rate constant (k) of 0.0026 ± 0.0016 and a corrected recovery (R) of 0.44 ± 0.17 after 17.6 min; which is 15% of the rate and 32% less recovery than that recorded for wild-type cells (k = 0.0174 ± 0.0113 and R = 0.65 ± 0.24 after 2 min; Figure 5; Table 3). The tub2-C354A mutant showed a more severe phenotype with a first-order rate constant (k) of 0.0014 ± 0.0009 and a corrected recovery (R) of 0.44 ± 0.23 after 18.6 min, only 8% of the rate and 32% less recovery than wild-type cells (Figure 5; Table 3). These results indicate that the tub2-C354 mutations substantially decrease the rate and total turnover of GFP-tubulin within the mitotic spindle. The reduced recovery in the mutants may reflect the limited duration of the analysis, even though it was 10 times longer than that needed for maximum wild-type recovery or a stable population of microtubule microtubules.

Spindles in our wild-type strain reached an average length of >8 μm before spindle separation and breakdown occurred. In the tub2-C354S mutant, the average spindle length was 5.1 ± 0.73 μm (n = 23, p < 0.001 vs. wild-type) at the time of spindle midzone separation and 17% of tub2-C354S spindles separated before reaching a length of 4.5 μm. The average length of a tub2-C354A mutant spindle at the time of spindle midzone separation was only 4.5 ± 0.98 μm (n = 20, p < 0.001 vs. wild-type, p = 0.016 vs. tub2-C354S), and 55% of the spindles separated before reaching 4.5 μm in length. In fact, 45% of tub2-C354A spindles separated when the spindle was between 3 and 4 μm long.

The rate of spindle elongation can be used to monitor the dynamics of polar microtubules (Straight et al., 1998; Severin et al., 2001). In budding yeast, spindle elongation during anaphase is biphasic (Kahana et al., 1995; Yeh et al., 1995; Straight et al., 1997). In our wild-type strain the initial elongation rate was 0.73 μm/min, dropping to 0.22 μm/min at a spindle length of 4 μm (Table 3). Many of the spindles in the mutant strains separated before reaching lengths greater than ~4.5 μm; however, a significant number reached lengths sufficient to allow us to determine the rate of elongation for spindles >4 μm long. The rates of spindle elon-
gation were significantly lower in the tub2-C354 mutants; however, both strains displayed biphasic spindle elongation kinetics (Table 3). The tub2-C354S spindles had rates of 0.27 and 0.16 μm/min for the fast and slow phases of anaphase B, respectively. For the tub2-C354A spindles these rates were 0.17 and 0.05 μm/min, respectively. These results indicate that suppression of microtubule dynamics reduces the overall rates of spindle elongation, thought to be driven by microtubule motors (Straight et al., 1998). However, the biphasic properties of spindle elongation are inherent to the process and not the polymer.

**In Vitro Dynamics of Wild-type and tub2p-C354S Microtubules**

To determine whether the effect of the tub2-C354 mutations on microtubule dynamics in vivo was solely attributable to the mutated tubulin molecule, we measured the in vitro dynamic parameters of individual microtubules formed from purified wild-type and tub2-C354S tubulins. We made use of His6-tag affinity chromatography to purify tubulin from each strain that was homogeneous by SDS-PAGE analysis (Figure 6). The data summarized in Table 4 show clearly that microtubules formed from tub2-C354S tubulin are much less dynamic than microtubules formed from wild-type tubulin. Sample microtubule lifetimes are presented in Figure 7. Both types of microtubules spent a very small amount of the total time in the shortening phase. Wild-type microtubules spent almost 100% of the time in the growth phase, whereas tub2p-C354S microtubules spent about half the time in the paused phase. In addition to this difference, the growth and shortening rates of the mutated microtubules were 33 and 6%, respectively, of the corresponding rates for wild-type microtubules. Dynamicity was reduced by 90%. The mutation also reduced the catastrophe frequency and increased the rescue frequency substantially. Wild-type microtubules experienced a catastrophe, on average, every 21.5 min. Tub2p-C354S microtubules experienced a catastrophe, on average, every 130 min (Table 4). Of the four catastrophes recorded for tub2p-C354S microtubules, three were followed by a rescue event. By contrast, wild-type microtubules showed no rescue events after 20 catastrophes. In Figure 7C the sample lifetimes are presented over a shorter time period and clearly show the differences in the catastrophes of wild-type and tub2p-C354S microtubules.

**Table 3. Spindle elongation rates and FRAP parameters in wild-type and tub2-C354 mutants**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Wild-type</th>
<th>tub2-C354S</th>
<th>tub2-C354A</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elongation rate (μm/min)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial (fast) phase*</td>
<td>0.73 ± 0.10 (7)</td>
<td>0.27 ± 0.09 (12)</td>
<td>0.17 ± 0.08 (13)</td>
</tr>
<tr>
<td>Sequential (slow) phase*</td>
<td>0.22 ± 0.08 (7)</td>
<td>0.16 ± 0.06 (8)</td>
<td>0.05 ± 0.01 (3)</td>
</tr>
<tr>
<td>FRAP parameters*</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First-order rate constant</td>
<td>0.0174 ± 0.0113 (9)</td>
<td>0.0026 ± 0.0016 (7)</td>
<td>0.0014 ± 0.0009 (7)</td>
</tr>
<tr>
<td>Half-time to recovery, s</td>
<td>52 ± 24 (9)</td>
<td>354 ± 186 (7)</td>
<td>600 ± 236 (7)</td>
</tr>
<tr>
<td>Corrected recovery (R)</td>
<td>0.65 ± 0.24 (8)</td>
<td>0.44 ± 0.17 (7)</td>
<td>0.44 ± 0.23 (7)</td>
</tr>
</tbody>
</table>

Results are reported as the mean ± 1 SD. Number of events is in parentheses.

* Elongation rates for spindles between 2 and 4 μm long.  
* Elongation rates for spindles greater than 4 μm long.  
* FRAP parameters were determined by analyzing an average recovery period after photobleaching of 2, 18, and 19 min for wild-type, tub2-C354S, and tub2-C354A spindles, respectively. The corrected recovery is the ratio of the fluorescence intensity of the bleached pole to the unbleached pole after recovery.

Figure 6. SDS-PAGE gels and Western blot of yeast tubulin. (A) Coomassie Blue–stained SDS-PAGE gel of purified tubulin from MGY1. Lane 1, 0.5 μg yeast tubulin standard; lane 2, 25 μg 100,000 × g supernatant; lane 3, 10 μg DE52 eluate; lane 4, 10 μg Ni-NTA pass-through; lane 5, 1 μg Ni-NTA eluate. (B) Western-blot of gel shown in A. Staining was with the anti-α-tubulin antibody clone B-5–1-2 and a horseradish peroxidase–conjugated anti-mouse antibody as the secondary antibody (Sigma-Aldrich, St. Louis, MO). Detection was accomplished with 4-chloro-1-napthol and H2O2. Because of its low concentration in the 100,000 × g supernatant, tubulin could not be detected in this fraction. (C) Coomassie Blue–stained SDS-PAGE gel loaded with 2 μg of purified tubulin from MGY1-C354S.
DISCUSSION

In tub2-C354 Mutants a Single Stable Cytoplasmic Microtubule Finds the Bud Site and Facilitates Spindle Positioning, but not Orientation

Wild-type budding yeast cells contain an average of three and as many as six cytoplasmic microtubules in unbudded cells (Shaw et al., 1997b; Tirnauer et al., 1999). In contrast, tub2-C354 mutant cells contained one stable and persistent cytoplasmic microtubule throughout the entire unbudded phase of cell growth. Microtubule localization to the bud is believed to involve a search and capture mechanism, which relies on microtubule dynamic instability to effectively probe the entire cell cortex (Carminati and Stearns, 1997). This dynamic property of microtubules has been shown to be an efficient mechanism for probing intracellular space (Holy and Leibler, 1994). Another mechanism of microtubule localization to the bud site has been proposed that involves directed-transport of microtubule ends along polarized actin filaments to the bud (Yin et al., 2000). In wild-type cells, cytoplasmic microtubules interact transiently with the future bud site and bud cortex, but the dynamic nature of the microtubules typically results in detachment and depolymerization out of the bud (Adames and Cooper, 2000; Beach et al., 2000). However, in all the tub2-C354 mutant cells examined, the cytoplasmic microtubule stopped moving and became stationary, with its end in the vicinity of imminent bud growth before bud emergence was visible. The tip of the tub2-C354 mutant microtubule remained associated with the emerging bud throughout bud growth. Thus, a minimally dynamic microtubule is able to locate the site of future bud growth without the aid of a dynamic instability-based microtubule probing mechanism. The minimum requirement to find the incipient bud site is simply a microtubule that is long enough to interact with the cell cortex. It would seem that the pivoting motion that the cytoplasmic microtubules undergo might be more important in finding the bud site than dynamics. Thus, at least in the tub2-C354 mutants, the directed-transport mechanism appears responsible for locating the microtubule structure at the bud site, possibly with the help of actin filaments (Theesfeld et al., 1999; Yin et al., 2000).

Defects in spindle position and orientation can be caused by changes in microtubule dynamics (Tirnauer et al., 1999; Kosco et al. 2001). In the tub2-C354 mutants, the spindle was positioned near the bud neck at the onset of anaphase. However, in the mutants the spindle was not oriented properly along the mother-bud axis in a larger percentage of the cells compared with wild-type cells. The single cytoplasmic microtubule was stable throughout G2/M and was ~1 µm in length during preanaphase. The failure of tub2-C354 mutants to maintain spindle orientation may reflect their inability to maintain cytoplasmic microtubule-cortical interactions at the distal ends of the mother cell and the bud.

Dynamics of Wild-type and tub2p-C354 Microtubules In Vivo and In Vitro

Our previous studies with the tub2-C354 mutants demonstrated that the mutations produced phenotypes that were consistent with an increase in microtubule stability (Gupta et al., 2001). To examine the effects of these mutations on microtubule stability in more detail, we analyzed microtubule dynamics in both wild-type and tub2-C354 mutant cells using a GFP-Tub1p fusion protein. The wild-type values we obtained for the microtubule dynamic parameters of growth rate, shrinkage rate, catastrophe frequency, and rescue frequency in wild-type cells were in agreement with those reported by others (Carminati and Stearns, 1997; Shaw et al., 1997b; Tirnauer et al., 1999; Adames and Cooper, 2000; Vogel et al., 2001; Kosco et al., 2001). However, all aspects of microtubule dynamics were drastically reduced in both tub2-C354 mutants. Consistent with our previous findings (Gupta et al., 2001), the tub2-C354A mutation decreased dynamics more than the tub2-C354S mutation. Depending on the phase of the cell cycle and the mutant, microtubule growth and shrinkage rates were reduced by 69–92%, catastrophe and rescue frequencies by 51–99%, and dynamics by 77–99%. In addition, in vivo results suggest that a threshold of microtubule dynamics is required for proper progression through the cell cycle. The more severe tub2-C354A mutant

Table 4. In vitro dynamics of wild-type and tub2-C354S microtubules

<table>
<thead>
<tr>
<th>Time spent in</th>
<th>Wild-type</th>
<th>tub2-C354S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Growth phase (%)</td>
<td>99.6</td>
<td>53.5</td>
</tr>
<tr>
<td>Shrinkage phase (%)</td>
<td>0.40</td>
<td>0.57</td>
</tr>
<tr>
<td>Attenuated phase (%)</td>
<td>0.00</td>
<td>46.0</td>
</tr>
<tr>
<td>Total time, min</td>
<td>43.1</td>
<td>520</td>
</tr>
<tr>
<td>Growth rate (µm/h)</td>
<td>10.6 ± 1.9 (32)</td>
<td>3.5 ± 1.2 (14)</td>
</tr>
<tr>
<td>Shrinkage rate (µm/min)</td>
<td>102.7 ± 18.4 (22)b</td>
<td>6.1 ± 2.1 (11)b</td>
</tr>
<tr>
<td>Catastrophe (min⁻¹)</td>
<td>0.0464 ± 0.010 (20)</td>
<td>0.0077 ± 0.004 (4)</td>
</tr>
<tr>
<td>Rescue (min⁻¹)</td>
<td>&lt;0.0023 (0)</td>
<td>0.0058 ± 0.003 (3)</td>
</tr>
<tr>
<td>Dynamicity (dimer/s)</td>
<td>16.6</td>
<td>1.9</td>
</tr>
</tbody>
</table>

a For wild-type dynamics 32 microtubules grown from 11 axonemes were analyzed. For tub2-C354S dynamics 16 microtubules grown from 8 axonemes were analyzed. Rates and frequencies are reported as the mean ± 1 SD. The number of events is in parentheses.
b Shrinkage events exceed catastrophes due to brief pauses during disassembly (see Figure 7C).

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Microtubule dynamics in vitro were decreased substantially by the tub2-C354S mutation. The changes in vitro dynamics parameters paralleled those observed in vivo. The one exception was the rescue frequency. The frequency of rescue in vivo was reduced by 50–98% by the tub2-C354S mutation. In vitro, however, rescue frequency was increased greater than 2.5-fold in the tub2-C354S microtubules. This apparent discrepancy can be explained on the basis of the stability of yeast microtubules. Wild-type microtubules depolymerize very rapidly in vitro, and rescue events are never observed (this study; Davis et al., 1993; Sage et al., 1995a, 1995b). However, the slow depolymerization rate for tub2-C354S microtubules in vitro enhanced the opportunity for rescue events, thus explaining the increase in rescue frequency. In contrast, the decreased number of microtubule catastrophes in the mutant cells reduced the opportunity for rescue to occur as well. The large decreases in the frequency of catastrophe and shrinkage rate in vivo and in vitro caused by substituting either a serine or alanine residue for cysteine 354 in Tub2p clearly illustrate the strong stabilizing effect of these mutations on microtubules. Because the in vitro studies were conducted with purified tubulin, the results demonstrate that the extreme microtubule stability is an intrinsic property of the mutated protein. The results further indicate that the intrinsic properties of microtubules are dominant relative to proteins that modulate dynamics in vivo.

Changes in Microtubule Dynamics During the Cell Cycle

The tub2-C354 mutants provide the most dramatic example to date of cell-cycle–specific modulation of microtubule dynamics in budding yeast. Microtubule dynamics varied as much as 6- to 8-fold during the cell cycle in these cells. The majority of microtubule dynamics modulation occurred because of changes in transition frequencies (up to 18-fold) and the proportion of time spent growing or shrinking (up to 10-fold in each case). These results indicate that microtubule dynamics can be altered to produce either net disassembly (small-budded) or net assembly (late-anaphase) of cytoplasmic microtubules. The decrease in length of the single microtubule in small-budded cells indicates that microtubule depolymerization is responsible for nuclear migration to the bud neck (Adames and Cooper, 2000; Beach et al., 2000).

In the wild-type cells, cytoplasmic microtubules became progressively less dynamic and more persistent as the cells approached mitosis. We found microtubule dynamics to be reduced 50% between G1 and anaphase (Table 1). This was achieved mainly by about a 25% reduction in growth and shrinkage rates and an increase in the proportion of time microtubules spent paused. The modulation of microtubule dynamics throughout the cell cycle in budding yeast indicates that cellular factors alter the intrinsic dynamic properties of microtubules.

Spindle Dynamics in tub2-C354 Mutant Cells

Spindle microtubule dynamics and the rate and extent of spindle elongation were reduced in the mutants. The results of the quantitative analysis in vivo correspond to the in vitro microtubule dynamics data, indicating that there is no nu-
clear or spindle-specific factor that can override the effects of the tubulin mutation. The anaphase spindles did not reach wild-type lengths (>8 μm), and the second, slower phase of spindle elongation was absent in many of the mutant cells. In the more severe tub2-C354A mutant, half of the spindles separated before reaching 4 μm in length. However, the spindle pole bodies segregated properly in all the mitotic events observed. Spindles that elongated significantly beyond 4 μm displayed the same type of biphasic elongation as observed in wild-type strains. In accordance with the less dynamic microtubules, the rates of spindle elongation were correspondingly lower in the tub2-C354A mutant relative to the tub2-C354S mutant. This finding indicates that the regulation of microtubule dynamics is important in determining the rate of spindle elongation. Previous evidence suggests that microtubule growth rate factors can control the rate of spindle elongation (Masuda and Candè, 1987). Experiments in yeast have shown that the bipolar nature of spindle elongation results from the balanced influence of various microtubule motor proteins (Straight et al., 1998). It is believed that kinesin-like motors influence spindle elongation by generating sliding forces on the interpolar microtubules in the spindle midzone. The results herein indicate that the bipolar rates are dependent on changes in the rate of polar microtubule elongation. The bipolar nature of the mutant spindle kinetics indicates that the balance of factors involved in spindle behavior contribute to the qualitative aspects of spindle elongation, but the restricted microtubule dynamics influences the quantitative rates of elongation. Clearly, the regulation of microtubule dynamics can directly influence microtubule-based processes such as spindle assembly and elongation.

The midanaphase checkpoint has been characterized as a pause in spindle elongation (at ~4 μm in length) in response to DNA damage (Yang et al., 1997), but the mechanism of the midanaphase checkpoint is not understood. Spindle elongation in the tub2-C354 mutants frequently paused at ~4 μm in length, suggesting possible involvement of the midanaphase checkpoint. However, the fact that the short tub2-C354 mutant spindles separated prematurely rather than remaining paused or elongating suggests an alternative explanation: that the difficulties observed in spindle elongation are structural and result from stabilized microtubules. In addition to the midanaphase checkpoint, the bipolar transition separates periods of spindle elongation that are controlled by different cellular factors, e.g., Cin8p and Kip1p (Straight et al., 1998), and may represent fundamentally different mechanisms of microtubule-based spindle elongation. The tub2-C354 mutant microtubules may be less capable of performing the functions associated with the second phase of spindle elongation.

C354 in β-Tubulin and Microtubule Dynamics

The dynamics of microtubules varies according to the species (Davis et al., 1993; Detrich et al., 2000) and the isotype of tubulin within a species (Panda et al., 1994; Gonçalves et al., 2001). Such variations are due to multiple differences in amino acid sequences. Our results demonstrate the sensitivity of microtubule dynamics to a single change in the primary sequence of tubulin, a point that has also been demonstrated with mutations of putative GTP binding site residues (Sage et al., 1995a, 1995b; Anders and Botstein, 2001; Dougherty et al., 2001). C354 in β-tubulin is situated at the αβ-dimer interface and is unlikely to be directly involved in longitudinal or lateral interactions in microtubule protofilaments. In our previous article we speculated that the mutation at C354 might produce a change in the structure of the tubulin dimer that could translate into stronger interprotofilament interactions in the microtubule (Gupta et al., 2001). This residue appears to be located at or near the colchicine binding site (Bai et al., 1996). It has been proposed that colchicine and other antimitotic drugs that modify microtubule dynamics mimic naturally occurring compounds that regulate microtubule function (Wilson and Jordan, 1995). The activities of microtubule regulatory agents that bind to the colchicine site may be mediated through contacts with C354 in β-tubulin.

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