Kinesin-5 Is Dispensable for Bipolar Spindle Formation and Elongation in Candida albicans, but Simultaneous Loss of Kinesin-14 Activity Is Lethal

Irsa Shoukat,a Corey Frazer,b John S. Allinghama

aDepartment of Biomedical and Molecular Sciences, Queen’s University, Kingston, Ontario, Canada
bDepartment of Molecular Microbiology and Immunology, Brown University, Providence, Rhode Island, USA

ABSTRACT Mitotic spindles assume a bipolar architecture through the concerted actions of microtubules, motors, and cross-linking proteins. In most eukaryotes, kinesin-5 motors are essential to this process, and cells will fail to form a bipolar spindle without kinesin-5 activity. Remarkably, inactivation of kinesin-14 motors can rescue this kinesin-5 deficiency by reestablishing the balance of antagonistic forces needed to drive spindle pole separation and spindle assembly. We show that the yeast form of the opportunistic fungus Candida albicans assembles bipolar spindles in the absence of its sole kinesin-5, CaKip1, even though this motor exhibits stereotypical cell-cycle-dependent localization patterns within the mitotic spindle. However, cells lacking CaKip1 function have shorter metaphase spindles and longer and more numerous astral microtubules. They also show defective hyphal development. Interestingly, a small population of CaKip1-deficient spindles break apart and reform two bipolar spindles in a single nucleus. These spindles then separate, dividing the nucleus, and then elongate simultaneously in the mother and bud or across the bud neck, resulting in multinucleate cells. These data suggest that kinesin-5-independent mechanisms drive assembly and elongation of the mitotic spindle in C. albicans and that CaKip1 is important for bipolar spindle integrity. We also found that simultaneous loss of kinesin-5 and kinesin-14 (CaKar3Cik1) activity is lethal. This implies a divergence from the antagonistic force paradigm that has been ascribed to these motors, which could be linked to the high mitotic error rate that C. albicans experiences and often exploits as a generator of diversity.

IMPORTANCE Candida albicans is one of the most prevalent fungal pathogens of humans and can infect a broad range of niches within its host. This organism frequently acquires resistance to antifungal agents through rapid generation of genetic diversity, with aneuploidy serving as a particularly important adaptive mechanism. This paper describes an investigation of the sole kinesin-5 in C. albicans, which is a major regulator of chromosome segregation. Contrary to other eukaryotes studied thus far, C. albicans does not require kinesin-5 function for bipolar spindle assembly or spindle elongation. Rather, this motor protein associates with the spindle throughout mitosis to maintain spindle integrity. Furthermore, kinesin-5 loss is synthetically lethal with loss of kinesin-14—canonically an opposing force producer to kinesin-5 in spindle assembly and anaphase. These results suggest a significant evolutionary rewiring of microtubule motor functions in the C. albicans mitotic spindle, which may have implications in the genetic instability of this pathogen.

KEYWORDS Candida albicans, yeast, kinesin, microtubule dynamics, mitotic spindle
The mitotic spindle is a highly dynamic microtubule (MT)-based structure that undergoes a distinct set of morphological changes in order to correctly attach, orient, and then separate sister chromatids in the dividing cell. Kinesin motor proteins play major roles in shaping and organizing MTs within the spindle over the course of cell division. Early in mitosis, evolutionarily conserved kinesin-5 proteins cross-link the overlapping plus ends of interpolar MTs from newly duplicated centrosomes (spindle pole bodies in yeast) and then slide them apart via plus-end-directed motility to establish spindle bipolarity (1–8). Genetic or chemical inhibition of kinesin-5 activity produces monopolar spindles or inward collapse of preanaphase spindles, usually leading to cell death (9–11). This spindle defect arises from loss of outward forces needed to counterbalance the inward forces supplied by MT minus-end-directed kinesin-14 motors, which pull spindle poles together (10, 12, 13). In many organisms, a nearly normal spindle phenotype can be restored by inactivating or depleting cells of kinesin-5 and kinesin-14 simultaneously because this force imbalance is eliminated (10, 13–16). In this experimental scenario, pushing forces generated by MT growth are sufficient to promote spindle pole separation and bipolar spindle assembly (17–20).

This interplay of motor and MT forces has been studied extensively in the model yeasts *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe* (10, 17, 18, 20–22). *S. cerevisiae* encodes two kinesin-5 homologs, Kip1 and Cin8, that have overlapping, but nonequivalent functions during mitosis (1, 2), while *S. pombe* encodes a single kinesin-5, named Cut7 (23). All three of these proteins form homotetramers that exhibit bidirectional motility, and all of them function in bipolar spindle assembly and cross-link parallel MTs to help focus kinetochore clusters (24–29). They are also important for stabilizing the overlapping array of MTs at the anaphase spindle midzone and for promoting and regulating timely anaphase spindle elongation (17, 22, 30–34). In both yeast species, loss or inhibition of kinesin-5 function is lethal. However, simultaneous inactivation of their kinesin-14 motors (Kar3Cik1 and Kar3Vik1 in *S. cerevisiae* or Pkl1 and Klp2 in *S. pombe*) neutralizes kinesin-5 deficiency (35–38), highlighting the importance of keeping inward and outward forces acting on the spindle in balance. In contrast to the lethality of kinesin-5 loss, bipolar spindles are able to form in the absence of kinesin-14 activity, but are either short and disorganized, or their MT minus ends are unfocused and extend past the opposite spindle pole (39, 40). Our studies of the homologous motors in the opportunistic fungus *Candida albicans* indicate that these phenotypes, and the opposing relationship of kinesin-5 and kinesin-14 proteins in spindle regulation, are not as highly conserved among eukaryotes as previously thought.

*C. albicans* is a close relative of *S. cerevisiae* and *S. pombe*, but it encodes only one kinesin-5 and one kinesin-14 motor, named CaKip1 and CaKar3, respectively, the latter of which forms a heterodimer with a noncatalytic kinesin-like protein, CaCik1 (41). *C. albicans* is viable without CaKip1 (42), and cells lacking CaKar3Cik1 activity often arrest with a monopolar spindle or two dissociated half-spindles (41). Through further investigation of these unconventional phenotypes, we found that CaKip1 is not needed for bipolar spindle assembly or nuclear division, even though it exhibits the same cell-cycle-dependent localization as its homologs in budding yeast. However, *kip1Δ/Δ* spindles are shorter and intermittently disassemble prior to cell division. When spindle disassembly occurs, two or more independent bipolar spindles emerge that either segregate between the mother and daughter cells or elongate across the bud neck. Each bring portions of the nucleus with them, which are further subdivided when the spindles undergo anaphase. Rather than neutralizing these kinesin-5 deficiencies, we found that simultaneous loss of kinesin-14 activity is lethal. These results imply that *C. albicans* Kip1 and Kar3Cik1 have mostly overlapping rather than antagonistic functions in bipolar spindle assembly and that their combined loss cannot be compensated for by MT polymerization forces or other spindle-associated factors.
RESULTS

Localization of C. albicans Kip1 mirrors other yeast kinesin-5s. Like many mitotic proteins, the localization and function of kinesin-5 motors changes throughout the cell cycle. During spindle assembly, S. cerevisiae Kip1 and Cin8, as well as S. pombe Cut7, are enriched at the minus ends of nuclear MTs, toward the spindle poles (26, 43, 44). Here, they are thought to capture MTs emanating from neighboring spindle pole bodies (SPBs) to establish antiparallel MT interactions and provide outward sliding forces to support SPB separation (7, 23–27, 29, 45). Persistence of kinesin-5 near spindle poles in metaphase has been attributed to their interaction with kinetochores, or to kinetochore MTs (kMTs), where they could cross-link parallel kMTs and regulate their assembly dynamics to help achieve chromosome congression (28, 46, 47). Upon anaphase onset, kinesin-5 motors relocate toward the plus ends of interpolar microtubules (ipMTs), which overlap in an antiparallel array in the spindle midzone. Here, their MT cross-linking and plus-end-directed motility help stabilize and elongate the spindle, fully separating the two opposing SPBs, leading to final chromosome segregation (17, 22, 30–34). Recent studies suggest that this cell-cycle-dependent redistribution of yeast kinesin-5s in the spindle is enabled by their capacity for bidirectional motility (24–27, 48, 49).

The discovery that C. albicans is viable without CaKip1 (42) suggests that its localization and/or function may be different from those of other yeast kinesin-5s. However, when we imaged fields of unsynchronized cells expressing green fluorescent protein (GFP)-labeled CaKip1 and mCherry-labeled tubulin (Tub2), we observed similar cell-cycle-dependent motor localization patterns within the mitotic spindle as seen in other yeasts. In small-budded early mitotic cells, CaKip1 localized near one end of monopolar spindles (in which SPBs are adjacent) (Fig. 1A, row 1) and was found at both poles after SPB separation and bipolar spindle assembly (Fig. 1A, row 2). In cells that were entering anaphase, CaKip1-GFP fluorescence was dispersed along the length of the spindle. In late anaphase cells, CaKip1 accumulated at the spindle midzone. The same localization patterns were seen when we imaged individual cells expressing CaKip1-mScarlet and Tub2-Neon over the course of mitosis by time-lapse microscopy, although photobleaching affected the ability to detect CaKip1 at later time points (Fig. 1B). To determine whether midzone clustering of CaKip1 requires overlapping arrays of antiparallel ipMTs in this region, we imaged CaKip1-GFP in fields of unsynchronized cells lacking kinesin-14 activity (cik1Δ/Δ). In other yeasts, kinesin-14 is important for organizing antiparallel ipMT interactions in the midzone so that kinesin-5 motors can properly cross-link and slide antiparallel spindle MTs (39, 40, 50). Without kinesin-14 activity, CaKip1 remains exclusively near the poles of bipolar spindles and one pole of dissociated half-spindles, presumably due to paucity of antiparallel ipMT overlaps (Fig. 1C).

C. albicans forms bipolar spindles without kinesin-5. To understand the role of CaKip1 in mitosis, we used PCR- and CRISPR-based gene targeting to generate two independent homozygous CaKIP1 deletion strains. Wary that CaKip1 could be essential for cell growth (51), we also engineered a conditional CaKIP1 gene expression strain using the tetracycline-regulatable (TR) promoter system, which enables tight repression of CaKIP1 in the presence of doxycycline (DOX) (52, 53). Transformants of each strain were screened by PCR to confirm the intended gene modification (data not shown). We further used transcriptome sequencing (RNA-seq) analysis to confirm absence of CaKIP1 expression in the gene deletion strains (see Table S1 in the supplemental material). The RNA-seq data showed that there were no changes in expression of any other molecular motors or MT-associated proteins (MAPs) to suggest the presence of compensatory mechanisms from such proteins.

In dilution spot assays, all CaKip1-depleted strains displayed modest sensitivity to higher temperature, but were otherwise viable (Fig. 2A). However, in liquid culture, cells lacking CaKip1 activity proliferated slower than wild-type cells and contained a mixture of blastoconidia and cells with long extensions resembling pseudohyphae.
 Upon further visual inspection and quantification of the \textit{kip1}Δ/Δ strain by microscopy, we observed this hyperpolarized morphology in approximately 30% of the cells (Fig. 2C). These elongated cells indicate a delay in cell cycle progression or a cell cycle arrest and could mask a slower proliferation rate on solid growth medium by giving \textit{kip1}Δ/Δ dilution spots a similar appearance to the wild type. We also found that loss of \textit{Ca}Kip1 affected filamentous growth under hypha-inducing conditions. \textit{Ca}Kip1-depleted colonies formed a smaller halo of invasive growth on Spider medium (Fig. 2D), and cells grown in serum produced shorter germ tubes and fewer septa (Fig. 2E and F).

When we added a wild-type copy of \textit{CaKIP1} back into the \textit{kip1}Δ/Δ strain at the native locus, normal cell growth rate and cell morphology were restored (Fig. 2B to D), confirming that these defects were a direct consequence of \textit{CaKIP1} loss.

Expecting that the slow-growth phenotype of \textit{Ca}Kip1-depleted cells was caused by errors in mitotic spindle assembly, we imaged fields of unsynchronized wild-type and \textit{kip1}Δ/Δ blastoconidia expressing Tub2-mCherry and Kip1-GFP (a component of the spindle pole body) and examined their spindle structures. We found that most of the spindles in budded \textit{kip1}Δ/Δ cells (92%) formed a stereotypical bipolar spindle structure (Fig. 3A). However, nearly twice as many \textit{kip1}Δ/Δ cells had metaphase spindles (53.5%) compared to the wild type (28.9%) (Fig. 3B), and the mean length of \textit{kip1}Δ/Δ metaphase spindles was significantly shorter (\textit{kip1}Δ/Δ, 0.68 ± 0.01 μm; wild type, 0.93 ± 0.01 μm) (Fig. 3C). When we tracked progression of the mitotic phases by time-lapse microscopy, we observed that \textit{kip1}Δ/Δ cells took an average of 117.1 ± 7.8 min to initiate anaphase after a spindle had formed, whereas wild-type cells

\begin{figure}[h]
\begin{center}
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\caption{\textit{Ca}Kip1 exhibits similar localization to other kinesin-5s during the cell cycle. (A) Images of wild-type cells expressing Tub2-mCherry and Kip1-GFP (strain CF338). Representative cells from different stages of mitosis were selected. (B) Individual frames from time-lapse microscopy of cells expressing GAL-Tub2-mNeon and Kip1-mScarlet (strain CF443). (C) Images of \textit{cik1}Δ/Δ cells expressing GAL1-Tub2-mCherry and Kip1-GFP (CF340). All cells were obtained from logarithmically growing, unsynchronized cultures in SDC-sucrose medium at 30°C. Scale bars, 5 μm.}
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required only 80.0 ± 5.2 min on average (Fig. 3D and E). In contrast, when kip1Δ/Δ spindles did eventually elongate, there was no difference in the duration of anaphase compared to the wild type (Fig. 3F). These data show that C. albicans is not solely dependent on kinesin-5 activity for bipolar spindle assembly or late anaphase spindle
elongation, but CaKip1 is important for timely separation of spindle poles after spindle assembly.

Interestingly, kip1Δ/Δ spindles had longer and more numerous astral MTs than the wild type (Fig. 4A to C). In many eukaryotes, including C. albicans, the plus ends of astral MTs strike the cell cortex, where they are captured by the minus-end-directed MT motor protein dynein (54–56). When this happens, dynein can draw the MT, and the attached SPB, toward the cortical contact site to facilitate proper spindle positioning, elongation, and/or migration (57, 58). In S. cerevisiae, this activity of dynein assists Cin8 and Kip1 in anaphase spindle elongation, and simultaneous loss of dynein and Cin8 activity is lethal (22, 31). We found the dyn1Δ/Δ strain to be nonviable in the presence of the CaKip1-specific inhibitor aminobenzothiazole (ABT), suggesting that CaKip1 and dynein also have overlapping functions in C. albicans (Fig. 4D). Perhaps the longer and more numerous astral MTs in the kip1Δ/Δ strain are an adaptation to CaKip1 loss that provides more opportunities for MT capture and pulling events by dynein, which could promote both anaphase spindle elongation and SPB separation during spindle assembly.

A subpopulation of kip1Δ/Δ cells have multiple spindles and show atypical cell cycle dynamics. Similar to previous findings by Chua et al. (42), we observed that a significant proportion (~12%) of kip1Δ/Δ blastoconidia and all kip1Δ/Δ cells with a hyperelongated morphology, contained multiple spindles (Fig. 5A and B). In some cases, monopolar and bipolar spindles were simultaneously visible within the same budding cell (Fig. 5A, row 3). To determine how these extra spindles formed, we collected time-lapse images of kip1Δ/Δ blastoconidia expressing Tub2-GFP. All of the multip spindle blastoconidia that we tracked (n = 20) formed two short “bars” of tubulin
fluorescence in the mother cell as the new bud began to emerge. Once a new bud formed, we observed two different multispindle configurations. In 65% of the cells we imaged, one of the tubulin structures traversed the bud neck, while the other remained in the mother compartment (Fig. 5C, row 2, t = 6 min). Each short fluorescent bar then elongated simultaneously, suggesting that they had formed distinct bipolar spindles. However, both spindles broke apart or disintegrated once the cell divided (Fig. 5C, row 3, t = 36 min; see Movie S1 in the supplemental material). This phenotype suggests that Cak1p activity is important for anaphase spindle stability. In a smaller cohort of cells (45%), one or both of the spindles elongated across the bud neck and appeared to complete anaphase (not shown). Both of these multispindle phenotypes were recapitulated in wild-type cells treated with ABT (Fig. 5D).

We were intrigued by this spindle defect because a subpopulation of wild-type *C. albicans* cells exposed to the antifungal agent fluconazole (FLC) display abnormal numbers of spindles as well. In the presence of FLC, DNA replication and nuclear division proceed ahead of bud emergence and completion of cytokinesis, respectively. Harrison et al. (59) showed that when this happens, some nuclei re-fuse or fail to separate due to mitotic collapse, forming tetraploid progeny with extra spindle components. Therefore, we next used time-lapse microscopy to track nucleolar segregation (using Nop1-mScarlet) in ABT-treated wild-type cells that formed multiple spindles. While the “no-drug” condition showed stereotypical nuclear and spindle dynamics that were well coordinated with bud emergence and growth (Fig. 6, rows 1 and 2; see Movie S2 in the supplemental material), ABT-treated cells contained one large patch of Nop1-mScarlet fluorescence and two separate bars of Tub2-GFP when the bud...
emerged (Fig. 6, row 4; see Movie S3 in the supplemental material). This indicates that a bipolar spindle had already formed and broken apart before bud evagination. As bud growth continued, the Nop1 patch divided and migrated with each spindle. When these spindles were segregated to the mother and daughter cell (cell 1, 65% of the cells...
imaged), two smaller nuclear fragments were visible in each compartment after spindle elongation (four Nop1-mScarlet patches in total, cell 1, \( t = 95 \text{ min} \)). However, within 20 min, each pair of Nop1 patches coalesced as a result of mitotic collapse. When the spindles elongated across the neck (cell 2, \( 45\% \) of the cells imaged), each spindle divided a Nop1-mScarlet patch into two different pieces, again resulting in four separate nuclear fragments (cell 2, \( t = 45 \text{ min} \)). Here, the two Nop1 foci in each compartment were produced from different anaphase spindles. As the time-lapse continued, these Nop1 foci appeared to merge into one (Fig. 6, cell 2, \( t = 75 \text{ min} \)). These results suggest that the extra spindles seen in \( \text{kip1}^{\Delta/\Delta} \) cells and ABT-treated wild-type cells could have formed in nuclei that experienced mitotic collapse or after merging of nuclear fragments from two distinct spindles that completed anaphase.

**Simultaneous loss of CaKip1 and CaKar3/Cik1 function is lethal.** In many of the eukaryotic systems, inactivation of kinesin-14 rescues the lethal spindle defects arising from inhibition or loss of kinesin-5 activity and spindles recover the ability to complete a relatively normal mitotic cycle (12–16, 37, 38, 60–65). This has been rationalized as a restoration of force balance in the spindle, where compensatory spindle forces are provided by MT polymerization and cross-linking proteins (17, 34, 66). In spite of repeated attempts, we were unable to obtain a \( \text{kip1}^{\Delta/\Delta} \text{kar3}^{\Delta/\Delta} \) strain by traditional methods (data not shown), suggesting they are synthetically lethal. To confirm this, we constructed a \( \text{KIP1}^{\Delta/\Delta} \text{KAR3}^{\Delta/\Delta} \) strain containing only one functional copy of \( \text{KAR3} \) that is under the control of the maltose-inducible Mal2 promoter. Indeed, when we deactivated the Mal2 promoter by culturing this strain on glucose (YPD [yeast extract-peptone-dextrose] medium), cell growth was arrested, demonstrating that simultaneous loss of CaKar3 and CaKip1 is lethal (Fig. 7A). We also observed that \( \text{kar3}^{\Delta/\Delta} \text{ckl}^{\Delta/\Delta} \) cells were not viable in the presence of ABT (Fig. 7B). To visualize events leading up cell death by loss of CaKip1 and CaKar3 function, we imaged Tub2-GFP fluorescence in \( \text{kar3}^{\Delta/\Delta} \) cells treated with 100 \( \mu \text{M ABT} \) every 15 min by time-lapse microscopy to avoid photobleaching. After 165 min of imaging, we observed a short anaphase spindle that did not elongate further. Within 3 h, spindle structures disappeared and cells showed no tubulin fluorescence (Fig. 7C). These results demonstrate that kinesin-5- and kinesin-14 have more functional overlap in \( C. \text{albicans} \) than in other organisms.

**DISCUSSION**

In nearly every type of eukaryotic system studied, kinesin-5 activity is needed to push newly duplicated centrosomes or SPBs apart to establish spindle bipolarity (1–4, 8, 23). Kinesin-5s also cross-link and bundle parallel and antiparallel spindle MTs and are...
the major providers for outward forces during anaphase spindle elongation (1–4, 7, 8, 23, 67). Without them, most cells exhibit mono-astral spindles and are nonviable (2, 9–11). In contrast, our genetic data demonstrate that the sole kinesin-5 gene in C. albicans is not essential in diploid cells. Perhaps the recent finding that CaKIP1 is an essential gene in a haploid isolate of C. albicans (51) is an indication that CaKIP1 mutants exhibit a form of ploidy-specific lethality, which is shared by other yeast genes involved in mitotic spindle stability (68). An alternative explanation for this discrepancy is that there are differences in protein expression between the haploid and diploid proteomes (69). Our results further show that kinesin-5 is dispensable for spindle assembly and anaphase spindle elongation in C. albicans. The only other organisms reported to complete mitosis without kinesin-5 activity are Caenorhabditis elegans and Dictyostelium discoideum (70–72). Although their mechanisms for kinesin-5-independent bipolar spindle assembly and elongation are not yet known, it has been suggested that cytoplasmic dynein-mediated astral MT pulling forces are involved. We propose that dynein may also fulfill these roles in C. albicans in the absence of kinesin-5 activity for several reasons. In earlier studies, C. albicans cells lacking the heavy chain of cytoplasmic dynein or the p150Glu-d subunit of dynactin exhibited spindle position, orientation, and elongation defects, and dramatically slowed nuclear dynamics (73, 74). In the filament-forming fungus Ustilago maydis, it was shown that stationary dynein motors capture and pull on the plus ends of astral MTs that emanate from SPBs, drawing the attached SPB toward the cortical contact site (75). In our studies, astral MTs were much longer and more numerous in kip1Δ/Δ cells, which could increase the frequency of these MT capture and dynein-mediated pulling events for SPB separation and spindle elongation. We also found dyn1Δ/Δ mutants to be nonviable in the presence of the inhibitor ABT, suggesting that CaKip1 and dynein have overlapping functions. Our future studies aim to better understand this putative intersection of dynein and CaKip1 functions. We will also investigate the alternative possibility that CaKip1 has a direct role in limiting the number and length of astral MTs, based on recent evidence that kinesin-5s can act as length-dependent MT depolymerases at kinetochores (46, 76, 77).

Although kip1Δ/Δ cells readily assembled metaphase spindles, these spindles were shorter and delayed in transitioning to anaphase relative to wild-type cells. This suggests a role for CaKip1 in maintenance of the bipolar spindle prior to anaphase, which is an important kinesin-5 function in other fungi and in Xenopus and Drosophila (3, 4, 45, 78). The redistribution of CaKip1-GFP fluorescence along the spindle in early anaphase supports such a role. The metaphase-to-anaphase delay in kip1Δ/Δ cells may also explain their lower growth rate in liquid cultures. Our observation that kip1Δ/Δ spindles sometimes broke apart and then reassembled two new bipolar spindles also supports this function and implies that CaKip1 is acting as a MT cross-linker within the spindle. An alternative explanation for these short, unstable spindles is that CaKip1 regulates kMT dynamics, which is important for congression of bioriented sister chromosomes in metaphase. Indeed, S. cerevisiae’s Cin8 is important for kinetochore clustering/positioning near the SPBs by cross-linking kMTs and promoting the disassembly of long kMTs (28, 46). In S. pombe, Cut7 is recruited to the kinetochores by a spindle assembly checkpoint (SAC) protein, Mad1, to promote chromosome gliding toward the spindle equator (29). Recent EM reconstructions of C. albicans KIP1/kip1Δ spindles show disorganized kMTs (76), suggesting that chromosomes are not properly congressed at the spindle equator during metaphase. Without proper chromosome congression, mitotic errors are more likely to occur. Perhaps the short bipolar spindles we observed in kip1Δ/Δ cells are indicative of attempts to correct erroneous kMT attachments (76), and spindle disassembly occurs when they are not corrected. If these cells initiate DNA replication and attempt mitosis again, this could explain the extra spindles and nucleoli observed in a subpopulation of C. albicans kip1Δ/Δ cells. Further work will be needed to uncover whether or not these defects in nuclear dynamics lead to an increase in the prevalence of aneuploid cells.

Surprisingly, CaKip1 loss did not extend the duration of anaphase relative to wild
type, even though CaKip1-GFP accumulated at the midzone of late anaphase spindles; a site where it could exert outward MT sliding forces for spindle elongation. This is unique from other fungi and Drosophila embryos, which rely on kinesin-5 to cross-link overlapping antiparallel MTs in the spindle midzone and drive anaphase B spindle elongation via plus-end-directed motility (6, 17, 22, 30–34, 63). While we suspect that dynein provides pulling forces on the spindle to assist in anaphase spindle elongation in the absence of CaKip1, it is also possible that other kinesins or MT cross-linking components within the spindle are involved. In fission yeast, kinesin-6 provides additional MT-sliding forces to kinesin-5 at the spindle midzone for anaphase spindle elongation and dynein is not involved (17, 66). Although C. albicans has no kinesin-6 homolog in its genome, it encodes five other kinesin-like proteins in addition to CaKip1. Therefore, we have begun to generate strains lacking different combinations of these proteins in order to identify new collaborative roles of kinesins in mitosis.

By simultaneously disrupting kinesin-5 and kinesin-14 activities, we found that C. albicans displays a puzzling exception to the widely regarded spindle force-balance model (15, 37, 62, 79). Rather than providing antagonistic spindle forces, CaKip1 and CaKar3Cik1 may cooperate to focus and stabilize parallel and antiparallel interactions in certain areas of the spindle. In this regard, loss of both kinesins may reduce the number of MT cross-linking factors to an intolerable level that cannot support cell viability. Our previous finding that CaKar3Cik1-depleted cells often arrest with two monopolar half-spindles that become pulled apart before assembling a bipolar spindle, supports this idea (41). Combined with CaKip1 loss, MTs may not be well tethered at SPBs or fail to focus kinetochores, resulting in disorganized spindle structures that quickly break down.

C. albicans is a close relative of the model yeasts S. pombe and S. cerevisiae but is also an opportunistic fungal pathogen. An assortment of fitness attributes promote its pathogenicity (80), most of which arise by rapid genetic diversification within a population in response to stressful growth conditions as a means of adaptation (81, 82). Research has shown that aneuploidy accounts for much of this diversity (83–85), and recent findings suggest that aneuploidies could be induced or enabled by altered activity of mitotic kinesin motors under stress (39, 49, 86–88). As specific aneuploidies can confer resistance to antifungal drugs through altered gene copy numbers, it could be advantageous for C. albicans cells to regulate mitotic kinesins as a way to control aneuploidy occurrence. We are currently conducting studies to delineate the putative contributions of C. albicans kinesins to mitotic defects of cells under stress and to identify stress-specific regulatory factors that change kinesin activity to promote aneuploidy.

MATERIALS AND METHODS

Genetic manipulations. A list of C. albicans strains used in this study is presented in Table 1. The oligonucleotides used in strain construction are listed in Table 2. Gene disruption of the C. albicans KIP1 open reading frame (Candida Genome Database, orf19.8331; NCBI Gene ID, 3645256) was conducted by PCR-based gene targeting and CRISPR-Cas9 methods (89). PCR amplification was used to generate disruption cassettes where a selectable marker was flanked by approximately 50 bp of C. albicans genomic sequence immediately 5’ and 3’ of the KIP1 coding region. Disruption of KIP1 in a wild-type strain (CF027) was conducted sequentially. First a kip1::LEU2 cassette was amplified from pSN40 (90) using primers P118 and P119 and transformed into strain CF027. Correct kip1::LEU2 cassette integration was confirmed using primer pairs P120/P13 and P121/P14 for the upstream and downstream junctions, respectively. To disrupt the second KIP1 allele, a kip1::HIS1 cassette was amplified from pSN52 (90) using primer pair P118/P119 and transformed to create strain CF311. Integration of the disruption cassette at the correct location was confirmed by PCR amplification across the junctions using primers P120/P11 and P121/P12 for the upstream and downstream regions, respectively. CRISPR-Cas9-mediated kip1 deletion was conducted as previously described (89) using the custom guide RNA (gRNA) primer P247 and double-stranded donor DNA formed using primers P248/P249 to create the strain CF429 (89). To regulate the expression of KIP1, the tetracycline-repressible transactivator, the teto promoter, and the NAT flipper cassette were PCR amplified from pLC605 (kindly provided by Leah Cowen) using primers P240/P241. The PCR-amplified product was transformed into the heterozygous KIP1 strain to create strain CF436. Correct integration at the KIP1 locus was verified using primer pairs P120 and P242.

To demonstrate that mutant phenotypes are solely a result of loss of KIP1, add-back strains were created to reintroduce a wild-type copy of each gene. The KIP1 gene (±1,000 bp upstream/downstream)
was cloned into pClp10-based integration plasmids bearing the ARG4\(^{+}\) selectable marker using primers P128/P129 (41, 91). The integration plasmid was digested at a unique restriction site (PmII) to add back to the endogenous KIP1 region into CF311 to create CF354. Confirmation of integration of the pClp10-ARG4\(^{+}\) vector was done using P121/P170.

A strain lacking both CakKir1 and CakKar3/Cik1 function was created by deleting both copies of KIP1 and one copy of KAR3 and by placing the remaining functional KAR3 copy under the control of a maltose promoter as follows: one copy of the KAR3 ORF was disrupted using kar3::HIS1\(^{+}\) knockout cassette amplified using primers P199/P200, transformed into CF027, and confirmed using primers P201/P11 and P202/P12 for the upstream and downstream junctions, respectively. The ARG4::MAL2-KAR3 cassette was amplified using primer pair P212/P213 from pFA-A-ARG4-MALp (92) and transformed in kar3::HIS1\(^{+}\) to create CF411 (not shown). Integration of the cassette was confirmed using primers P201 and P16. KIP1 was disrupted using LEU2 (described above) and the SAT1 nourseothricin resistance marker to create the strain CF396. Correct kip1::SAT1 integration was confirmed with primer pairs P128/P17 and P129/P18.

Fluorescence tagging of KIP1\(^{+}\) in wild-type cells was accomplished using the method described by Gerami-Nejad et al. (93) and using long-tailed primers P137 and P187 and the plasmid pGFP-SAT1 as a template to create an integration cassette bearing approximately 50 bp of KIP1\(^{+}\) ORF immediately before the stop codon and of sequence 3' to the ORF. This cassette was transformed into the wild type (CF027) to create the KIP1::GFP-SAT1\(^{+}\) strain. Correct integration was confirmed by PCR using the primer pair P69 and P169. The same integration cassette was also transformed in cik1\(\Delta\) to create strain CF308 (not shown). KIP1-mScarlet was amplified using pScarlet plasmid PR8897 kindly provided by Richard Bennett using primers P284 and P285 and transformed into CF421 (pGAL1-Tub2-mNeon) to create CF443. Integration was confirmed using primers P254 and P121.

Strains expressing fluorescently labeled β-tubulin were constructed using the plasmids pGAL1-Tub2-GFP-SatR; pGAL1-Tub2-mCherry-Arg4::NEUTSL, or pGAL1-Tub2-mNeon-Arg4::NEUTSL using the previously described method (41) and adapted to further include sequence of the neutral NEUTSL locus, which was linearized using the restriction enzyme KpnI. pGAL1-Tub2-GFP was transformed into CF027 and CF311 to create CF289 and CF226, respectively. pGAL1-Tub2-mCherry was transformed into the wild-type, kip1\(\Delta\), and cik1\(\Delta\) strains to create CF363, CF368, and CF340, respectively. pGAL1-Tub2-mCherry was transformed into KIP1-GFP to create strain CF338. Correct integration for pGAL1 vectors was confirmed by PCR using primers P16 (or P17, depending on SAT/ARG markers) and P107. Induction of the GAL1 promoter, leading to expression of fluorescently tagged tubulin, was done by growing the cells in SDC-sucrose medium supplemented with 1% galactose. To visualize the nucleus, the nucleolar protein Nop1 was fluorescently labeled using pScarlet. The integration cassette was PCR amplified using primers P243 and P244 and transformed into CF289 to create CF417. Correct integration was confirmed by PCR using primers P254 and P246. To visualize spindle pole body structures, strains expressing SPC98-GFP were constructed as previously described (41).
## TABLE 2 Oligonucleotide primers used in strain construction

| Primer | Description | Sequence (5′ to 3′) |
|--------|-------------|---------------------|
| P118   | Long homologous tail knockout primer | GTTGGTTGTTTGTATTCTTTTCATCATTGGTTGTTTGGATTATACTATGATA |
|        | KIP1::HIS1/LEU2/ARG4 5′ | GACGACTTACTATCTACCATAGTTCAGGATTTTAGCTGTTGTTGTTGTTGTT |
| P119   | Long homologous tail knockout primer | AAAAAAACCTACAAATTTAAAACGATGGAGAACAAATGAGAT |
|        | KIP1::HIS1/LEU2/ARG4 3′ | ATATTTGCTTTATTTTATACTCATAGCATGTTGTTGTTGTTGTTGTT |
| P120   | −500 bp KIP1 check 5′ | CGCACAAGACCTTGGGCAACAGGAAAG |
| P121   | +500 bp KIP1 check 3′ | ATGGGCGCAATGGATACATG |
| P11    | HIS1 check right 3′ | AACAAGACTGCAATCATGTTG |
| P12    | HIS1 check left 5′ | ATTAGATACTTGTTGTGTCAGTT |
| P13    | LEU2 check left 5′ | AGATTCCCAACATTTGTGTC |
| P14    | LEU2 check right 5′ | AACATCTGACCCGTCG |
| P247   | KIP1 gRNA for fragment B stitching | CTTGAAAACATTAAAATTCTATGATTGTTATGGTTGTTGTTGTTGTTGTTGTT |
| P248   | KIP1 donor DNA with mini-AT 5′ | ATTTCTTCTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTTGTT |
| P249   | KIP1 donor DNA with mini-AT 3′ | ACAATTATATAACATGTGAGAAGA |

*The portion of primer homologous to plasmid template is in boldface and italic. The restriction enzyme cut site is underlined. Lowercase letters represent the mini-ADD-TAG sequence (mAT [gg]) for subsequent CRISPR-mediated gene editing (89).*
C. albicans transformation. Disruption cassettes, fluorescent tags, and complementation plasmids were transformed into C. albicans using the lithium acetate–polyethylene glycol (PEG) heat shock method as previously described with minor modifications (94). Incubation of cells with transforming DNA in lithium acetate-PEG solution was carried out for 2 h at 30°C with rotation. Heat shock was conducted at 43°C for 30 min. Transformations involving selection using the SAT1 gene were accompanied by a 4-h incubation in YPD (1% yeast extract, 2% peptone and 2% glucose) at 30°C to allow expression of the ClonNAT resistance gene before plating on selection medium.

C. albicans cell culture and growth assays. Strains were maintained on YPD plates. YPD was supplemented with 200 µg/ml nourseothricin (clonNat; Werner BioAgents) for selection of positive SAT1 gene integration. Selection for auxotrophic markers was conducted using synthetic dropout (SD) medium containing 0.66% yeast nitrogen base, 0.2% yeast dropout mix lacking uracil, arginine, leucine, and histidine, 2% glucose, and 200 mg/liter uridine and supplemented with 200 µg/liter histidine, leucine, and/or arginine where required. Experimental cultures were grown to mid-logarithmic phase in completely supplemented dropout medium (SDC) unless otherwise indicated. In order to assess the discovery rate was set at 95%. Cell pellets were flash frozen in liquid nitrogen, and genomic DNA-free total RNA was extracted from each pellet by grinding the fungal mass to a fine powder and resuspending it in 1 ml TRIzol (Ambion) solution and using the RNeasy mini-spin columns (Qiagen) following the manufacturer’s protocol. RNA quantification was carried out spectrophotometrically at 260 nm and 280 nm, and RNA integrity was evaluated by NanoDrop2000 (Thermo Scientific). Total RNA (1 µg/sample) was shipped to the National Research Council of Canada, DNA Sequencing Technologies Facility (Saskatoon, Canada), where further quality check was performed using a BioAnalyzer followed by short cDNA fragment synthesis using the TruSeq Stranded RNA-LT kit, and finally sequenced on an Illumina HiSeq 2500 platform according to the manufacturer’s guidelines (Illumina, USA). The DESeq2-based SARTools (v1.5.1) pipeline as previously described (93) was adopted for differential analysis of mapped C. albicans Assembly 22 RNA-seq count data. A BH P value adjustment was performed (96, 97), and the false-discovery rate was set at P < 0.05.

Data availability. The RNA-seq data that support the findings of this study are provided in Table S1 and are available at the Sequence Read Archive (SRA) under BioProject accession no. PRJNA579546.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00610-19.

TABLE S1, XLSX file, 0.1 MB.
MOVIE S1, AVI file, 3 MB.
MOVIE S2, AVI file, 0.1 MB.
MOVIE S3, AVI file, 0.3 MB.

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REFERENCES

1. Hoyt MA, He L, Loo KK, Saunders WS. 1992. Two Saccharomyces cerevisiae kinesin-related gene products required for mitotic spindle assembly. J Cell Biol 118:109–120. https://doi.org/10.1083/jcb.118.1.109.

2. Roof DM, Meluh PB, Rose MD. 1992. Kinesin-related proteins required for assembly of the mitotic spindle. J Cell Biol 118:95–108. https://doi.org/10.1083/jcb.118.1.95.

3. Sawin KE, LeGuellec K, Philippe M, Mitchison TJ. 1992. Mitotic spindle organization by a plus-end-directed, microtubule motor. Nature 359:540–543. https://doi.org/10.1038/359540a0.

4. Heck MM, Pereira A, Pesavento P, Yannoni Y, Spradling AC, Goldstein LS. 1993. The kinesin-like protein KLP61F is essential for mitosis in Drosophila. J Cell Biol 123:665–679. https://doi.org/10.1083/jcb.123.3.665.

5. Kashina AS, Baskin RJ, Cole DG, Wedaman KP, Saxton WM, Scholey JM. 1996. Opposing motor activities are required for the organization of microtubule bundles in the mammalian mitotic spindle. J Cell Biol 135:399–414. https://doi.org/10.1083/jcb.135.2.399.

6. Sharp DJ, McDonald KL, Brown HM, Matthies HJ, Walczak C, Vale RD, Sullivan W, Scholey JM. 1995. Phosphorylation by p34cdc2 regulates spindle association of human Eg5, a kinesin-related motor essential for bipolar spindle formation in vivo. Cell 83:1159–1169. https://doi.org/10.1016/0092-8674(95)90142-6.

7. Enos AP, Morris NR. 1990. Mutation of a gene that encodes a kinesin-like protein blocks nuclear division in Aspergillus nidulans. Cell 60:1019–1027. https://doi.org/10.1016/0092-8674(90)90330-n.

8. Saunders WS, Hoyt MA. 1992. Kinesin-related proteins required for structural integrity of the mitotic spindle. J Cell Biol 118:95–108. https://doi.org/10.1083/jcb.118.1.95.

9. Saunders WS, Koshland D, Eshel D, Gibbons IR, Hoyt MA. 1995. Saccharomyces cerevisiae kinesin- and dynein-related proteins required for anaphase chromosome segregation. J Cell Biol 126:301–313. https://doi.org/10.1083/jcb.126.2.301.

10. Saunders WS, Hoyt MA. 1992. Two Saccharomyces cerevisiae kinesin-related proteins required for microtubule cross-linking in fission yeast. Mol Biol Cell 3:1475–356. https://doi.org/10.1083/jcb.147.2.351.

11. Tao L, Mogliner A, Civelekoglu-Shoehly G, Wollman R, Evans J, Stahlberg H, Scholey JM. 2006. A homotetrameric kinesin-5, KLP61F, bundles microtubules and antagonizes Ndc in mitoty assays. Curr Biol 16:2392–2398. https://doi.org/10.1016/j.cub.2006.09.064.

12. Rincon SA, Lamson A, Blackwell R, Syrovatkina V, Frasier V, Paoletti A, Betterton MD, Tran PT. 2017. Kinesin-5-independent mitotic spindle assembly requires the antiparallel microtubule crosslinker Asel in fission yeast. Nat Commun 8:15286. https://doi.org/10.1038/s41467-018-06118-w.

13. Yukawa M, Yamada Y, Toda T. 2019.Suppressor analysis uncovers that MAPs and microtubule dynamics balance with the Cut7/kinesin-5 motor for mitotic spindle assembly in Schizosaccharomyces pombe. G3 (Bethesda) 9:269–280. https://doi.org/10.1038/s43386-018-00896.

14. Toso A, Winter JR, Garrod AJ, Amaro AC, Meraldi P, McNinch AD. 2009. Kinetochore-generated pushing forces separate centromeres during bi-polar spindle assembly. J Cell Biol 184:365–372. https://doi.org/10.1083/jcb.200809055.

15. Yukawa M, Kawakami T, Okazaki M, Kume K, Tang NH, Toda T. 2017. A microtubule polymerase cooperates with the kinesin-6 motor and a microtubule cross-linker to promote bipolar spindle assembly in the absence of kinesin-5 and kinesin-14 in fission yeast. Mol Biol Cell 28:3647–3659. https://doi.org/10.1091/mbc.E17-06-0361.

16. Geiser JR, Schott EJ, Kingsbury TJ, Cole NB, Totis LJ, Bhattacharyya G, Hoyt MA. 1997. Saccharomyces cerevisiae genes required in the absence of the Cin8-encoded spindle motor act in functionally diverse mitotic pathways. Mol Biol Cell 8:1035–1050. https://doi.org/10.1091/mbc.8.6.1035.

17. Saunders WS, Koshland D, Eshel D, Gibbons IR, Hoyt MA. 1995. Saccharomyces cerevisiae kinesin- and dynein-related proteins required for anaphase chromosome segregation. J Cell Biol 126:301–313. https://doi.org/10.1083/jcb.126.2.301.

18. Saavedra A, Winter JR, Garrod AJ, Amaro AC, Meraldi P, McNinch AD. 2009. Kinetochore-generated pushing forces separate centromeres during bi-polar spindle assembly. J Cell Biol 184:365–372. https://doi.org/10.1083/jcb.200809055.

19. Toso A, Winter JR, Garrod AJ, Amaro AC, Meraldi P, McNinch AD. 2009. Kinetochore-generated pushing forces separate centromeres during bi-polar spindle assembly. J Cell Biol 184:365–372. https://doi.org/10.1083/jcb.200809055.
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76. McCoy KM, Tubman ES, Claas A, Tank D, Clancy SA, O’Toole ET, Berman J, Odde DJ. 2015. Physical limits on kinesin-5-mediated chromosome congression in the smallest mitotic spindles. Mol Biol Cell 26:3999–4014. https://doi.org/10.1091/mbc.E14-10-1454.

77. Tubman E, He Y, Hays TS, Odde DJ. 2018. Kinesin-5-mediated chromosome congression in insect cell spindles. Cell Mol Bioeng 11:25–36. https://doi.org/10.1007/s12195-017-0500-0.

78. Barton NR, Pereira AJ, Goldstein LS. 1995. Motor activity and mitotic spindle localization of the Drosophila kinesin-like protein KLP61F. Mol Biol Cell 6:1563–1574. https://doi.org/10.1091/mbc.6.6.1563.

79. Cytrynbaum EN, Scholey JM, Mogilner A. 2003. A force balance model of early spindle pole separation in Drosophila embryos. Biophys J 84:757–769. https://doi.org/10.1016/S0006-3495(03)74895-4.

80. Brown AJ, Budge S, Kalori D, Tillmann A, Jacobsen MD, Yin Z, Ene IV, Bohovych I, Sandai D, Kastora S, Potrykus J, Ballou ER, Childers DS, Shahana S, Leach MD. 2015. Stress adaptation in a pathogenic fungus. J Exp Biol 217:144–155. https://doi.org/10.1242/jeb.088930.

81. Selmecki A, Forche A, Berman J. 2010. Genomic plasticity of the human fungal pathogen *Candida albicans*. Eukaryot Cell 9:991–1008. https://doi.org/10.1128/EC.00060-10.

82. Berman J. 2016. Ploidy plasticity: a rapid and reversible strategy for adaptation to stress. FEMS Yeast Res 16:flow020. https://doi.org/10.1093/femsyr/flow020.

83. Forche A, Abbey D, Pisithkul T, Weinzierl MA, Ringstrom T, Bruck D, Selmecki A, Forche A, Berman J. 2006. Aneuploidy and isochromosome acquisition of aneuploidy provides increased fitness during the evolution of antifungal drug resistance. PLoS Genet 5:e1000705. https://doi.org/10.1371/journal.pgen.1000705.

84. Mary H, Fouchard J, Gay G, Reyes C, Gauthier T, Gruget C, Pecreau J, Tournier S, Gachet Y. 2015. Fission yeast kinesin-8 controls chromosome congression independently of oscillations. J Cell Sci 128:3720–3730. https://doi.org/10.1242/jcs.160465.

85. Chen S, Stout JR, Dharmaiah S, Yde S, Calvi BR, Walczak CE. 2016. Transient endoreduplication down-regulates the kinesin-14 HSET and contributes to genomic instability. Mol Biol Cell 27:2911–2923. https://doi.org/10.1091/mbc.E16-03-0159.

86. Mary H, Fouchard J, Gay G, Reyes C, Gauthier T, Gruget C, Pecreau J, Tournier S, Gachet Y. 2015. Fission yeast kinesin-8 controls chromosome congression independently of oscillations. J Cell Sci 128:3720–3730. https://doi.org/10.1242/jcs.160465.

87. van Ree JH, Nam HJ, Jeganathan KB, Kanakkanthara A, de Vries J. 2016. Pten regulates spindle pole movement through Dlg1-mediated recruitment of Eg5 to centrosomes. Nat Cell Biol 18:814–821. https://doi.org/10.1038/ncb3369.

88. van Ree JH, Nam HJ, Jeganathan KB, Kanakkanthara A, de Vries J. 2016. Pten regulates spindle pole movement through Dlg1-mediated recruitment of Eg5 to centrosomes. Nat Cell Biol 18:814–821. https://doi.org/10.1038/ncb3369.

89. Nguyen N, Quail MMF, Hemday AD. 2017. An efficient, rapid, and recyclable system for CRISPR-mediated genome editing in Candida albicans. mSphere 2:e00149-17. https://doi.org/10.1128/mSphere Direc-10.1128/mSphereDirect.00149-17.

90. Noble SM, Johnson AD. 2005. Strains and strategies for large-scale gene deletion studies of the diploid human fungal pathogen Candida albicans. Eukaryot Cell 4:298–309. https://doi.org/10.1128/EC.4.2.298-309.2005.

91. Gola S, Martin R, Walther A, Dunkler A, Wendland J. 2003. New modules for PCR-based gene targeting in Candida albicans: rapid and efficient gene targeting using 100 bp of flanking homology region. Yeast 20:1339–1347. https://doi.org/10.1002/yea.1044.

92. Gerami-Nejad M, Berman J, Gale CA. 2001. Cassettes for PCR-mediated construction of green, yellow, and cyan fluorescent protein fusions in Candida albicans. Yeast 18:859–864. https://doi.org/10.1002/yea.738.

93. Walther A, Wendland J. 2003. An improved transformation protocol for the human fungal pathogen Candida albicans. Curr Genet 42:339–343. https://doi.org/10.1007/s00294-002-0349-0.

94. Varet H, Brillat-Guigue L, Coppee JY, Dillies MA. 2016. SARTools: a DESeq2- and EdgeR-based R pipeline for comprehensive differential analysis of RNA-Seq data. PLoS One 11:e0157022. https://doi.org/10.1371/journal.pone.0157022.

95. Benjamini Y, Hochberg Y. 1995. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J R Stat Soc Ser B Stat Methodol 57:289–300. https://doi.org/10.1111/j.2517-6161.1995.tb02031.x.

96. Benjamini Y, Yekutieli D. 2001. The control of the false discovery rate in multiple testing under dependency. Ann Stat 29:1165–1188. https://doi.org/10.1214/aos/1013699998.

97. Benjamini Y, Yekutieli D. 2001. The control of the false discovery rate in multiple testing under dependency. Ann Stat 29:1165–1188. https://doi.org/10.1214/aos/1013699998.

98. Sherwood RK, Bennett RJ. 2008. Microtubule motor protein Kar3 is required for normal mitotic division and morphogenesis in Candida albicans. Eukaryot Cell 7:1460–1474. https://doi.org/10.1128/EC.00138-08.