Spindle pole body-anchored Kar3 drives the nucleus along microtubules from another nucleus in preparation for nuclear fusion during yeast karyogamy

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Nuclear migration during yeast karyogamy, termed nuclear congression, is required to initiate nuclear fusion. Congression involves a specific regulation of the microtubule minus end-directed kinesin-14 motor Kar3 and a rearrangement of the cytoplasmic microtubule attachment sites at the spindle pole bodies (SPBs). However, how these elements interact to produce the forces necessary for nuclear migration is less clear. We used electron tomography, molecular genetics, quantitative imaging, and first principles modeling to investigate how cytoplasmic microtubules are organized during nuclear congression. We found that Kar3, with the help of its light chain, Cik1, is anchored during mating to the SPB component Spc72 that also serves as a nucleator and anchor for microtubules via their minus ends. Moreover, we show that no direct microtubule–microtubule interactions are required for nuclear migration. Instead, SPB-anchored Kar3 exerts the necessary pulling forces laterally on microtubules emanating from the SPB of the mating partner nucleus. Therefore, a twofold symmetrical application of the core principle that drives nuclear migration in higher cells is used in yeast to drive nuclei toward each other before nuclear fusion.

[Keywords: nuclear migration; microtubule; kinesin; karyogamy; spindle pole body; Kar3]

Supplemental material is available for this article.

Received September 19, 2012; revised version accepted December 27, 2012.
by targeting it to cytoplasmic MTs and SPBs [Meluh and Rose 1990; Page et al. 1994].

After mating is initiated by cell–cell fusion, the haploid nuclei of the mating cells become transported toward each other in a process termed nuclear congression [Rose 1996]. This involves specific regulation of MTs at the yeast centrosome, the spindle pole body (SPB). This mating-specific function of the SPB is initiated by mating signaling processes and leads to reorganization of the cytoplasmic MTs. Thereby, the γ-tubulin complex-binding protein Spc72, along with associated MTs, becomes released from its anchor protein, Nud1, at the outer plaque structure of the SPB and subsequently reappears at the half bridge appendage of the SPB. Here, it directly binds to the half bridge component Kar1, a tail-anchored protein that is essential for mating and SPB duplication in vegetative cells [Rose and Fink 1987; Spang et al. 1995; Pereira et al. 1999]. Upon establishment of cell–cell fusion at the site of cell polarity, the tip of the so-called shmoo, the half bridge-anchored MTs promote the migration of the two SPBs toward each other and thus of the entire nuclei. This process depends on the activity of the Kar3 motor protein.

To explain the mechanism of force generation for nuclear congression, two different models were proposed. Both models involve direct interactions between MTs from the opposite SPBs of the mating partners, however, with each having a different role for Kar3 [Molk and Bloom 2006]. The “sliding cross-bridge” model [Rose 1996] proposed overlapping anti-parallel MTs that slide along each other and drive the nuclei toward each other via their SPB-anchored minus ends. In this model, Kar3 acts as cross-linker for anti-parallel MTs and provides by its minus end-directed motility the pulling forces to drag the two SPBs together [Polaina and Conde 1982, Meluh and Rose 1990; Endow et al. 1994]. In addition, Kar3 was suggested to trigger shortening of opposing MTs at the SPBs [Rose 1996]. An alternative model proposed direct interaction of MTs at their plus ends and their subsequent depolymerization. This is proposed to lead to a shortening of the MTs and, by this, drive nuclear congression. In this model, the Kar3/Cik1 heterodimer would function as a linker of the MT plus ends and induce depolymerization of the MTs from their plus ends, thus promoting the congression movements [Molk et al. 2006].

Upon completion of congression, nuclear fusion occurs in three steps in which each of the two bilayers of the nuclear membranes and, subsequently, also the SPBs fuse with each other [Melloy et al. 2007], giving rise to one nucleus containing a diploid genome and one so-called fusion SPB [Byers and Goetsch 1975]. This process requires a set of additional Kar proteins for the membrane and SPB-related fusion processes [Melloy et al. 2009].

In this study, we used electron tomography, live-cell imaging, and simulation to quantitatively analyze the functional dynamics of MT interactions, MT anchor proteins, and the kinesin-14 Kar3 at high resolution. Our data suggest a new model for nuclear congression in which Kar3 is part of a module consisting of its light chain Cik1 and the γ-tubulin complex-binding protein Spc72, anchored to the nucleus via the half bridge component Kar1 of the yeast SPB. We show that this module underlies congression-specific MT organization. It anchors MT minus ends and, via Kar3 associated with the mating-specific Cik1 light chain, exerts the forces necessary for congression on MTs emanating from the SPB of the mating partner’s nucleus. This permits the nuclei to move toward each other without the need of direct MT–MT interactions. Therefore, the Kar3-mediated migration of nuclei along MTs in yeast mating is analogous to nuclear migration processes in higher cells.

Results

MT organization during nuclear congression

Previous genetics and microscopic analyses led to the proposal of two models for how MTs could function to promote nuclear congression [Rose 1996; Molk et al. 2006]. Hoping to distinguish between them, we imaged cells using electron tomography because its resolution in the range of 5–10 nm provides a precise snapshot of all MTs in the cell. We reconstructed wild-type cells that have undergone cell–cell fusion but in which the two nuclei were still separated, one on each side of the cell fusion pore [n = 14 tomograms] [three zygotes shown in Fig. 1A; four other zygotes are shown in Supplemental Fig. S1A]. In the segmented three-dimensional (3D) models, we observed distances between the two SPBs of the mating cells in the range of 0.16–2.57 μm. These values exhibited a good anticorrelation with the area of the openings between the cells [Supplemental Fig. S1B], indicating that the 3D models represent cells at different stages of congression, from early to late. Notably, inspection of the nuclear morphology identified 11 zygotes that exhibited protrusion-like nuclear deformations pointing toward the mating partner, each of which contained an SPB at the tip [Fig. 1A]. We detected one to six MTs nucleated per SPB [3.12 ± 1.24 SD]. The origin of each MT was clearly indicated by the capped MT minus ends, which were found in close proximity [less than ~15 nm] to the half bridge of each SPB [Fig. 1B]. This is consistent with the previous observation of MTs emanating from the half bridge upon pheromone stimulation in yeast [Byers and Goetsch 1975]. The plus ends, characterized by their open morphology, were detected in the cytoplasmic space between and around the two nuclei.

The MT organization in the 14 zygotes, however, was not consistent with any of the published models because direct MT–MT interactions in the space between the two nuclei were dramatically underrepresented. Instead, MTs of variable length, with several of them much longer than the distance between the two SPBs, were observed. We thus scanned along MTs originating clearly from one SPB to identify whether they would come in close proximity to other subcellular structures, including opposite MTs.

In all 14 zygotes, we identified eight cases in which one MT plus end appeared close to the lattice of an opposite MT [<100 nm], but none of these MTs exhibited an extended anti-parallel alignment. In six out of these eight...
zygotes with MT–MT adjacencies, the nucleus was strongly deformed, with the SPB located at the tip of a protrusion that extended toward the other nucleus. Furthermore, we observed 11 zygotes with one to four long MTs (two on average) that passed the opposite SPB in close proximity (Fig. 1C; Supplemental Fig. S1C; Supplemental Movie S1), with a median lateral MT–SPB distance of 58.8 nm ($n=21$) and a maximum distance of 106 nm (Fig. 1D). Interestingly, these 11 zygotes were also the ones harboring nuclear protrusions, whereas the three remaining zygotes, which showed no protrusions, also clearly did not show MTs in close proximity of opposing SPBs.

Assuming that the nuclear protrusions constitute a signature for pulling forces applied by a MT to an SPB, the co-occurrence of protrusions and MT–SPB adjacencies could point to the SPBs as the sites that connect to MTs from opposite SPBs. In contrast, only six out of the 11 zygotes with protrusions, but also two of the three non-protruding zygotes, exhibited MT–MT adjacencies. Altogether, our observations suggest a new model for nuclear congression, in which no direct MT–MT interactions, but rather SPB–MT interactions, are necessary to generate forces during nuclear congression. In such a model, the forces pulling the nuclei together could be generated, for example, by a minus end-directed MT motor that is anchored on the outer surface of the SPB. The observed nuclear protrusions would also be a direct signature of the forces.

Figure 1. MT organization during nuclear congression reveals long MTs in the vicinity of opposite SPBs. (A) Mating populations of wild-type [WT] yeast cells were cryofixed, freeze-substituted, and plastic-embedded. Electron tomograms of 300-nm sections were acquired from mating pairs. The panels show snapshots of the 3D-rendered situations of early, middle, or late stages (from top to bottom) of nuclear congression (Supplemental Movie S1), judged by the distances between opposite SPBs. (Light blue) Plasma membrane; (purple) nuclear envelope; (yellow) SPBs; (green) MTs. Bar, 300 nm. (B) Electron tomographic slice and the corresponding area in the 3D model showing a SPB in a mating cell with half bridge-anchored MTs. Bar, 100 nm. (C) Close-up view of four SPB models showing long MTs from the opposite SPB in their vicinity. Bar, 30 nm. The distances were measured between these long MTs and opposite SPBs. (D) Distances between long MTs and SPBs. Median, upper, and lower quartiles [box] are shown.
Long MTs from one SPB connect the opposite SPB during nuclear congression

The observed median distance of 58.8 nm between a MT and its opposite SPB is comparable with the size of a kinesin motor protein (Howard 1996). The minus end-directed kinesin Kar3 is known to be essential for nuclear congression (Meluh and Rose 1990; Endow et al. 1994), and by using immunofluorescence microscopy in pheromone-stimulated cells, Kar3 was found to localize to MTs and their plus tips as well as to SPBs (Meluh and Rose 1990). To investigate the role of Kar3 during nuclear congression in living cells, we generated an endogenously expressed and fully functional N-terminally tagged GFP-Kar3 fusion protein (Supplemental Fig. S2A). GFP-Kar3 localized to the SPBs labeled with Spc72-3mCherry throughout nuclear congression. In addition, GFP Kar3 also localized to the lattice of MTs and at MT plus ends, in agreement with previous data from immunofluorescence microscopy (Fig. 2A; Supplemental Movie S2; Meluh and Rose 1990). Consistent with the observations made by electron tomography (Fig. 1), we also detected long MTs that appear to extend across the gap between the two nuclei beyond the opposing SPBs (Supplemental Fig. S2B). This shows that long MTs nucleated at one SPB pass in close proximity and beyond the opposite SPB. Therefore, Kar3, localized at the opposite SPB, could act laterally on these MTs to generate the pulling forces necessary for nuclear congression. Furthermore, we quantified the dynamics of nuclear congression using Spc42-GFP-labeled SPBs. Cytoplasmic TagBFP expressed in one mating partner was used as a proxy for cell–cell fusion [Fig. 2B]. The plots of the distance between the SPBs as a function of time after cell–cell fusion (n = 30) (Fig. 2C) revealed three distinct phases of nuclear congression. In phase i, the SPBs move little, while in phase ii, they approach each other with a speed of 1.27 ± 0.35 μm/min. In 11 out of 30 traces, no clear phase i could be observed, and the SPBs seemed to migrate almost immediately after cell–cell fusion. Finally, in phase iii, the SPBs remain close to each other at a distance of <1 μm for a variable period of time (22 out of 30 cells). Completion of nuclear congression, indicated by a merge of the two SPB signals, was detected in 24 out of the 30 cells investigated within the
15 min of the image acquisition periods. This last phase may correspond to fusion of the nuclear membranes, followed by fusion of the nucleoplasms and finally of the SPBs [see also Fig. 5A in Melloy et al. 2007]. Interestingly, the measured speed of congression matched the velocity of Kar3 [1–2 μm/min] measured in vitro [Endow et al. 1994; Chu et al. 2005].

Altogether, the results from electron tomography and live-cell imaging argue for a role of long MTs in promoting nuclear congression through interactions with the opposite SPB, possibly via Kar3.

A cytoplasmic SPB-anchored pool of Kar3 promotes efficient nuclear congression

The kinesin Kar3 is present at the SPB and along MTs. To distinguish the function of these two different pools of the motor protein during nuclear congression, we depleted Kar3 from the MT lattice only. To do so, we fused GFP-Kar3 directly to the C terminus of the essential SPB outer plaque component Cnm67 [Muller et al. 2005]. This alteration did not cause growth defects in cells still containing a wild-type copy of the KAR3 gene, indicating that Cnm67 remained functional and that the Cnm67-GFP-Kar3 fusion protein does not exhibit any dominant-negative effects in vegetative cells. Moreover, the Cnm67-GFP-Kar3 protein fusion did not rescue the slow-growth defect of a kar3Δ mutant, indicating that it does not complement the mitotic nuclear function of Kar3 [Supplemental Fig. S2C]. We analyzed the intensity of the GFP-Kar3 signal on SPBs and along MTs using pheromone-treated CNM67-GFP-KAR3 kar3Δ cells. The Cnm67-GFP-Kar3 fusion protein was found to localize at the SPB, but in contrast to GFP-Kar3 cells, no signal was detected along MTs and in the cytoplasm [Fig. 2D, Supplemental Fig. S2D]. Using DAPI staining to score nuclear fusion in mating cells, we found that the Cnm67-GFP-Kar3 fusion was able to restore the mating defect of kar3Δ cells to ~40% of the one observed for wild-type cells [Supplemental Fig. S2E].

We measured SPB congression dynamics in these cells [Fig. 2E, Supplemental Movie S3]. In contrast to wild-type cells, we observed that all CNM67-GFP-KAR3 kar3Δ mating cells exhibited a clearly detectable phase i of up to 200 sec [Fig. 2F]. Subsequent nuclear congression in phase ii, however, occurred with wild-type speeds [1.45 ± 0.42, n = 25, P > 0.05], but in the majority of the observed cases in the SPBs of the CNM67-GFP-KAR3 kar3Δ cells, no clearly detectable phase iii was observed [16 out of 25] [Fig. 2F]. Instead, the signals of the two SPBs directly merged with each other.

From these results, it appears that Kar3, when forced to localize to the outer plaque of the SPB, is able to exert motor protein activity sufficient to promote nuclear congression with wild-type speeds. However, initiation and termination of congression seem to occur in an altered manner. This may be caused by the absence of Kar3 at the MT lattice as well as the altered localization of Kar3 to the outer plaque instead of the half bridge of the SPB.

Cytoplasmic Kar3 localizes to the SPB through interaction with Spc72

Our results so far suggest that Kar3 is the likely candidate for the interaction of SPBs with cytoplasmic MTs from opposite SPBs. Interestingly, Kar3 localization to the SPB in pheromone-stimulated cells was shown to be dependent on Kar1 [Vallen et al. 1992], implying that Kar3 would localize at the cytoplasmic side of the SPB. However, in mitotic cells, Kar3 was found on the nuclear side of the SPB [Zeng et al. 1999]. Therefore, to confirm that Kar3 indeed localizes at the cytoplasmic side of the SPB during wild-type mating, we used pheromone-stimulated cells and measured the position of the GFP-Kar3 signal relative to the SPB central plaque marker Spe42-mCherry using light microscopy. Gaussian fitting was used to detect the origin of the GFP-Kar3 fluorescence with subpixel accuracy. Moreover, Spc72-GFP and Spc110-GFP, which are known to localize at the cytoplasmic and nuclear sides of the SPB, respectively, were used to validate this approach [Supplemental Fig. S2F,G]. Indeed, we found that Kar3 localizes at the cytoplasmic side of the SPB during mating [Fig. 3A].

We next used fluorescence recovery after photobleaching (FRAP) to investigate Kar3 dynamics at the SPB and obtained a mean residence time of GFP-Kar3 at the SPB of 15 sec [Fig. 3B]. Kinesin-14 minus end-directed and non-processive motor proteins, such as Kar3, bind transiently to MTs with a typical residence time <200 msec [Chen et al. 2011]. The much longer residence time of Kar3 at the SPB thus likely reflects that the localization mechanism is independent of the binding activity of Kar3 to MTs. This was already suggested by the SPB localization of a truncated Kar3 fragment lacking the motor domain in pheromone-stimulated cells in the presence of nocodazole [Meluh and Rose 1990]. To identify a possible molecular anchor of Kar3 at the cytoplasmic side of the SPB, we screened for two-hybrid interactions of Kar3 with known SPB components [Elliott et al. 1999]. We detected an interaction between Kar3 and the C-terminal coiled-coil domain of Spc72 but not with the N-terminal domain [Fig. 3C] through which Spc72 binds to Spc97 and Spc98 of the γ-tubulin complex. This latter interaction of Spc72 underlies the nucleation and SPB anchorage of MT minus ends [Knop and Schiebel 1997]. In vegetative cells, Spc72 binds to Nud1 at the SPB outer plaque, whereas it binds to Kar1 at the half bridge during mating [Pereira et al. 1999; Grueneberg et al. 2000]. Also, Cik1, the Kar3 light chain involved in mating and mitosis [Page and Snyder 1992], was found to interact with Spc72 [Fig. 3C]. Using kar3Δ, cik1Δ, and vik1Δ deletion mutants for two-hybrid testing, we observed a mutual dependency of Cik1 and Kar3 for their interaction with Spc72, whereas Vik1 was not required [Fig. 3C]. This is fully consistent with the observation that Cik1 localization to SPBs in pheromone-treated cells depends on Kar3 [Benanti et al. 2009], and the localization of Kar3 to the cytoplasmic side of the SPB depends on Cik1 [Fig. 3D], as also previously suggested by immunofluorescence analysis [Page et al. 1994].
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These results indicate a dual role of Spc72 at the half bridge of the SPB for the interaction with MTs during nuclear congression. Spc72 binds MT minus ends via the γ-tubulin complex [Pereira et al. 1999] and simultaneously also anchors the Kar3/Cik1 complex.

Kar1-anchored Spc72/Kar3 machinery is sufficient to promote MT minus end movements

The SPB half bridge component Kar1 is anchored with its C-terminal membrane domain in the nuclear envelope [Spang et al. 1995], whereas a region in the N-terminal half (amino acids 116–236) interacts directly with Spc72 [Pereira et al. 1999]. kar1 mutations in the Spc72 interaction domain, such as kar1Δ13, kar1Δ15, or kar1-1, are the only ones known that lead to a unilateral mating defect (Conde and Fink 1976; Rose and Fink 1987; Vallen et al. 1992, Pereira et al. 1999) in which it is sufficient to have the mutation in only one of the two mating partners to abrogate mating completely [to the level observed when both partners contain the mutation]. Our finding that Spc72 binds Kar3/Cik1 in addition to MT minus end-binding proteins is consistent with this. Presumably, such kar1 mutants do lose simultaneously both types of SPB-MT interactions. Thereby, MTs nucleated and anchored at the wild-type KAR1 SPB fail to interact with the kar1 SPB, which itself is also unable to anchor and nucleate MTs.

To test this model further, we used live-cell microscopy. Imaging GFP-Kar3 in kar1Δ15 cells, we indeed observed the loss of cytoplasmic, but not nuclear, GFP-Kar3 signal from the SPB, along with the detachment of Spc72 and the anchored MTs, upon stimulation with pheromone [Fig. 4A–C; Supplemental Movie S4]. This is consistent with previous observations showing that a motor-less domain of Kar3-lacZ fusion protein does not bind to kar1Δ13 SPBs in pheromone-stimulated cells [Vallen et al. 1992].

To investigate the nuclear morphology and the MT organization as a function of Spc72 binding to the half bridge, we generated tomograms from zygotes of kar1Δ15 mated to wild-type cells [n = 8] (Supplemental Movie S5). This always revealed one SPB completely free of MTs nucleated from the opposite—and presumably wild-type—SPB. Furthermore, nuclear protrusions were absent in these zygotes, suggesting that their formation depends on Kar1-anchored Spc72 and functioning nuclear congression [Fig. 4D]. In addition, we observed free MTs in the cytoplasm [n = 7 in a total of 8 tomograms], as reported for pheromone-stimulated cells [Rose and Fink 1987]. These MTs each exhibited a capped end, most likely the minus ends that detached from the kar1Δ15 SPBs [Fig. 4D]. Strikingly, most of the capped minus ends were detected in close proximity (<100 nm) to the lattice of a MT from the opposite SPB [n = 6; median distance of 30.5 nm] (Fig. 4E,F).

This result could suggest active capturing of these minus ends by other MTs. To investigate this possibility, we followed the fate of detached MTs in kar1Δ15 strains using live-cell imaging and GFP-tubulin and Spc72-mCherry as labels. In several cases, we observed capturing of a free end of a released MT by a MT of the opposite SPB [n = 20 out of 22] [Fig. 4G,H]. In all recorded events, the captured end was labeled with Spc72, and in the majority of the cases [n = 17 out of 22], subsequent movement toward the minus end of the capturing MT [as indicated by the position of the opposite SPB] [Supplemental Movie S6] was observed. In some cases, we could follow their movement until they...
Figure 4. Kar1-anchored Spc72/Kar3 provides MT minus end-directed mobility. (A) Release of cytoplasmic MT together with a minus end-attached GFP-Kar3 pool from the SPB (followed with mCherry-labeled Spc42) in a pheromone-stimulated kar1Δ15 cell (Supplemental Movie S4). The nucleus is indicated with the dashed line that limits the GFP-Kar3 nuclear signal, the arrow points to the detached MT minus end, and the plus sign (+) identifies the plus end at the shmoo tip. Bar, 2 µm. (B) Released cytoplasmic MT harboring a minus end-attached GFP-Kar3 [left] and Spc72-3mCherry [right] pools detached together from the SPB in a pheromone-stimulated kar1Δ15 cell. The nucleus is indicated with the dashed line that outlines the GFP-Kar3 nuclear signal, the arrow points to the detached MT minus end, and the plus sign (+) identifies the plus end at the shmoo tip. Bar, 2 µm. (C) SPB-retained GFP-Kar3 in pheromone-stimulated kar1Δ15 cells with detached cytoplasmic MTs localizes to the nuclear side of the SPB (24 cells, n = 26, labeled Nuc). Experimental settings are as in Figure 3A. (D) 3D models of mating between MATα kar1Δ15 and MATα wild-type (WT) partners. Typically, one SPB was found to be free of cytoplasmic MTs (kar1Δ15 SPB), and released MTs with free capped minus ends are visible in the cytoplasm (Supplemental Movie S5). (Light blue) Plasma membrane; (purple) nuclear envelope; (yellow) SPBs; (green) MTs. Bar, 300 nm. (Panels i,ii) Electron tomographic slices show the region around the indicated areas in the model to the left. Bar, 30 nm. (Panels 1–4) 3D models show close-up views of different kar1Δ15 SPBs. Bar, 30 nm. (E) 3D models of mating events between kar1Δ15 and wild-type cells. (Red circles) The minus ends of detached MTs in the vicinity of another MT; (purple) nuclear envelope; (yellow) SPBs; (green) MTs. Bar, 300 nm. (Right) Distances between detached MT minus ends and the lattice of the closest MT. Median, upper, and lower quartiles (box) are shown. Individual outliers (cross) are shown if above/below quartile + 1.5 × (interquartile range) (whiskers). (F) Frames from a movie showing the capture and alignment of a detached MT in a kar1Δ15 to a MT from the opposite wild-type SPB. GFP-Tub1 [left] and Spc72-3mCherry [right] were detected (Supplemental Movie S6). (H) Behavior of Spc72 dots during mating in wild-type and mutant cells. Spc72-3mCherry- and GFP-tubulin-labeled cells of both mating types [with mating type-specific genotypes, as indicated] were mixed to allow mating to occur followed by movie recording in cells undergoing mating. The behavior of Spc72-3mCherry was quantified with respect to detachment from one of the two SPBs [presumably the kar1Δ15 SPB] and its position and movement were quantified with respect to cytoplasmic MTs that emanate from the SPB with attached Spc72.
reached a parallel alignment with the capturing MT. The observed minus end-directed movement suggests the presence of active Kar3 motor protein, together with Spc72, at the minus ends of the released MTs. Indeed, in a kar3Δ background, none of the Spc72-containing MTs showed minus end-directed movements along other MTs [Fig. 4H]. These results indicate that the detached Kar3/Spc72 module at MT minus ends in kar1Δ15 cells is fully functional, thereby implying that Kar3 bound to Spc72 at the half bridge in wild-type cells is active and able to promote minus end-directed movements.

**An asymmetric MT organization is sufficient for nuclear congression**

It was previously reported that the overexpression of Spc72 in kar1-1 mutant cells partially rescued mating with wild-type cells [Fig. 5A; Pereira et al. 1999]. In vegetative cells, overexpression of Spc72 led to the formation of a large aggregate of the protein at the SPB outer plaque [Souës and Adams 1998] and an increased number of cytoplasmic MTs [Knop and Schiebel 1998]. Whereas Spc72 detaches from the SPB in kar1Δ15 mutants stimulated with pheromone [Pereira et al. 1999], we observed persistent binding in the case in which Spc72 was overexpressed, along with attached MTs [Supplemental Fig. S3A]. This could explain how Spc72 overexpression in kar1Δ15 cells can rescue the mating defect. Interestingly, the mating defect was even more efficiently rescued when Spc72 was overexpressed in the wild-type instead of the kar1Δ15 mating partner [Fig. 5A; Supplemental Fig. S3B–D]. We used electron tomography to investigate this further. We observed a significant increase of the MT number at the wild-type SPB (n = 3) [Fig. 5B], all of which appeared to be nucleated from aggregates of Spc72 at the SPB outer plaque [Fig. 5C]. In contrast, we did not observe MTs nucleated from the kar1Δ15 SPB [Fig. 5D]. Nonetheless, we noticed in one tomogram a cell in an advanced state of nuclear congression, as indicated by the close proximity of the two SPBs from facing nuclei, in spite of the absence of MTs at one SPB [Fig. 5D, panel ii]. In this tomogram, the nucleus containing the anchored MTs exhibited a characteristic nuclear protrusion otherwise found only in wild-type tomograms but never in kar1Δ15 zygotes outside the overexpression context. This seemingly contradicted the idea of Spc72 being the sole anchor at the SPB for the machinery that mediates lateral MT interactions.

To further investigate how Spc72 overexpression in the wild-type partner of a wild type × kar1Δ15 mating could rescue mating with an asymmetric MT organization, we analyzed MT dynamics in mating cells. To distinguish the genotypes of the mating partners, we labeled Spc72 with 3mCherry. Before cell–cell fusion, the overexpressing cell showed nuclear congression that appeared to progress with MTs nucleated only at the SPB of the Spc72-overexpressing cell [Fig. 6A; Supplemental Movie S7]. In addition, however, we observed that small but detectable amounts of Spc72 localized at the SPB of the kar1Δ15 cell after cell–cell fusion, while no MTs were found to emanate from this SPB [Fig. 6B]. It is therefore likely that Spc72 at the...
In silico simulation of nuclear congression

Altogether, our data suggest a model in which the minus end-directed motor Kar3 anchored at the cytoplasmic side of the SPB promotes nuclear congression by pulling the SPBs along the MTs of the mating partner. To test whether this is physically plausible, we simulated the process within accurate 3D geometry using the Cytosim platform [http://www.cytosim.org] (Nedelec and Foethke 2007). The simulation is based on molecular descriptions of MT dynamics, the interactions of the Kar3/Cik1 dimer with MTs via its motor and nonmotor domain, and the SPB via its nonmotor domain [Fig. 7A,B]. To complete the list of parameters required for the simulation [see Supplemental Table S1], we measured the number of Kar3 motors at the SPB in pheromone-stimulated cells. The GFP-Kar3 fluorescence at the SPB in mating wild-type cells was quantified and calibrated using the fluorescence of signal of the kinetochore component Nuf2-GFP as a reference [Joglekar et al. 2006; Lawrimore et al. 2011]. This estimated an average of 143 ± 69 GFP-Kar3 at the SPB [mean ± SD, n = 34] (data not shown).

We simulated the system for 20 min after cell-cell fusion [Supplemental Movie S8] and extracted the trajectories of the SPBs as a function of time.

Three models were simulated: Model 1 contained Kar3 only at the SPBs, model 2 contained Kar3 at the SPBs and the plus ends of MTs, and model 3 contained only MT plus end-localized Kar3 (Fig. 7A). We observed robust nuclear congression for simulations of models 1 and 2 [Fig. 7C]. For model 3, the simulation did not lead to nuclear congression. A comparison of the trajectories from the simulations with wild-type live-cell imaging data [Fig. 2B] identified the best similarity of these data with the results obtained from model 2, where Kar3 localizes to SPBs, and a moderate number (three to five)
of freely diffusible Kar3/Cik1 is found at the plus ends of MTs (Fig. 7C; Supplemental Movie S9). Model 1 yielded, with the same set of dynamic parameters, trajectories in agreement with the live-cell imaging data obtained from CNM67-GFP-KAR3 kar3Δ cells that do not contain MT-localized Kar3, including a prolonged phase i (Figs. 2F, 7D). Therefore, the results from models 1 and 2 could point to a function of MT-localized Kar3 that promotes a shortening of phase i, possibly by orienting MTs toward the other SPB. In this model, we also tested the effect of two different amounts of Kar3 at the SPB and found that this only marginally influences the length of phase i and had no impact on the speed of congression (Fig. 7D).

We next analyzed the organization of MTs in the different models with respect to the localization of the plus ends of the MTs. In the absence of Kar3 at the MTs [model 1], we detected, on average, 30% of the cells at any given time point where a MT plus end was close to a MT of the opposite SPB. In model 2, we found that this value exhibited a peak with 64.4% of such cells 30–50 sec after cell–cell fusion. Later on, this percentage returned back to a value of 30% as observed in model 1. This indicates that the observed shortening of phase i in model 2 when compared with model 1 may be due to MT plus end-localized Kar3 motors interacting with the lattice of MTs originating from the opposite SPB. The frequency of such events in the simulation is furthermore consistent with the observation of such events in eight of the 14 tomograms [Fig. 1; Supplemental Fig. S1].

Phase iii, corresponding to a pausing phase prior to final SPB and nuclear fusion, was not covered by any of the simulations. This is due to our simplified description of MT anchorage at the SPB and the missing information about SPB substructures and membrane-related events that occur in the last steps of nuclear congression and fusion.

We convolved the simulation data to simulate live-cell imaging of cytoplasmic MTs and make the model directly comparable with experimental data [Supplemental Fig. S4]. This led to images highly reminiscent of the ones observed previously [Molk et al. 2006], in which MT–MT overlap zones between the SPBs are absent. Thus, our model is also consistent with these data, although it differs from the model published together with them.

In conclusion, the stochastic simulations using physics-based description of the molecular processes accurately reproduced the first two congression phases. The results suggest that phase i corresponds to the time needed for MTs to find their partner SPB, and phase ii starts when Kar3 engages with MTs from the opposite SPB. Nuclear congression phase iii, corresponding to a pausing phase prior to final SPB and nuclear fusion, was not covered by any of the simulations. This is due to our simplified description of MT anchorage at the SPB and the missing information about SPB substructures and membrane-related events that occur in the last steps of nuclear congression and fusion.

We convolved the simulation data to simulate live-cell imaging of cytoplasmic MTs and make the model directly comparable with experimental data [Supplemental Fig. S4]. This led to images highly reminiscent of the ones observed previously [Molk et al. 2006], in which MT–MT overlap zones between the SPBs are absent. Thus, our model is also consistent with these data, although it differs from the model published together with them.

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Figure 7. Physics-based simulation of nuclear congression using Cytosim. (A) Cartoon of the three different models used to simulate nuclear congression. Green circles indicate the possible localizations of Kar3 in the different models. (B) Snapshots of a simulation (model 1) 1 sec after cell fusion. In the simulations, the MTs can grow [green arrowhead], undergo catastrophe, and shrink [red arrowhead]. (C) Plots showing the distance between the two SPBs in mating cells from experiments and simulations. Thick solid lines show the median distances, and shaded areas indicate the upper and lower quartiles. The average of 100 simulations each is shown for each model. In model 1, Kar3 motors localize only at the SPB (Supplemental Movie S8). In model 2, the motors are present at the SPBs and on MT plus ends (green) (Supplemental Movie S9). For MT-localized motors, this model also contained, in addition to the SPB-bound motors, 2000 freely diffusing Kar3/Cik1 complexes with a binding activity to growing MT plus ends and the MT lattice via the nonmotor and motor domains. This resulted in 4.67 ± 1.65 motors per growing MT. Model 3 corresponds to model 2 without SPB-bound Kar3/Cik1 complexes (red) (Supplemental Movie S10). For more information and parameters, see the Materials and Methods and Supplemental Table S1. Experimental data show wild-type (WT) SPB congression (black; from Fig. 2C). (D) Results for model 1 simulations containing different numbers of SPB-bound motors [140 (blue solid line) and 280 motors per SPB (blue dashed line)], and experimental data show CNM67-GFP-KAR3 kar3Δ SPBs congression (black; from Fig. 2F).
congression then proceeds with a constant speed determined by the speed of Kar3. Our results show that MTs interacting with motors anchored at the opposite SPB are indeed a valid scenario for nuclear congression. We identified that plus end-localized Kar3/Cik1 can shorten the first search phase, but these interactions appear to be not sufficient to promote congression.

Discussion

The MT machinery is specifically regulated to perform nuclear congression. While it was well established that Kar3 together with Cik1 act on cytoplasmic MTs to drive nuclear migration, the limited resolution of light microscopy left several possibilities open for the underlying functional organization. The two pre-existing models (Rose 1996; Molk et al. 2006), based on the reported localization of Kar3 on MTs and/or at MT plus ends, involved direct MT–MT interactions. These models also satisfied the asymmetric mating defect of kar1 mutants caused by detachment of the MTs from one of the SPBs during mating (Rose and Fink 1987; Vallen et al. 1992; Pereira et al. 1999). Our initial observation showing that no clear MT–MT interactions could be detected was in apparent conflict with this kar1 phenotype, but this was resolved with the discovery that the Kar3/Cik1 localization to the SPB is dependent on an interaction with Spc72. Thus, Kar1 binding of Spc72 in mating cells recruits a functional bipartite MT-organizing module for MT nucleation and minus end binding via interaction of the N-terminal domain with the γ-tubulin complex (Knop and Schiebel 1998) and via binding to Kar3/Cik1 [Fig. 3] to move on MTs from the opposite SPB. This suggests a simple model in which a mating-specific module of Kar1/Spc72/Kar3/Cik1 functionalizes the half bridge of the SPB to take a lead in congression. The nucleus, which is connected to the SPB via the nuclear membrane and the attached chromosomes, follows this movement up to the point at which two SPBs of the mating partner encounter each other and nuclear fusion is initiated.

Involved in multiple cellular functions at different developmental stages, Kar3 illustrates how the motor protein can be used differently by regulating the expression of associated proteins. In its different functions, Kar3 is associated with Cik1 or Vtk1, thus forming distinct complexes (Manning et al. 1999). Transcriptional regulation of Cik1 into a cytoplasmic form upon stimulation of the mating process [Benanti et al. 2009] appears to be sufficient to adjust the localization and thus the functional activity of the motor. In mitotic spindles, Kar3 localizes on the nuclear side of the SPB [Zeng et al. 1999] in a Vtk1-dependent manner and in a Cik1-dependent manner on the plus ends of the nuclear interpolar MTs [Gardner et al. 2008]. During mating, Vtk1 is not expressed, and Kar3 interacts only with Cik1 [Manning et al. 1999]. In shmooing cells, cytoplasmic Kar3/Cik1 has been proposed to maintain depolymerizing MT plus ends at the shmoo tip (Meluh and Rose 1990; Maddox et al. 2003). This is likely due to an interaction of Kar3 with the mating-specific Ga subunit of a trimeric G protein involved in pheromone signaling [Zaichick et al. 2009]. Pheromone stimulation increases the expression of KAR3 and CIK1 via activation of the transcription factors Ste12 and Kar4 [Kurihara et al. 1996; Gammie et al. 1999; Lahav et al. 2007] and underlies the expression of the shorter NLS-less cytoplasmic Cik1 variant [Manning et al. 1999; Benanti et al. 2009]. Thus, the cytoplasmic localization of Cik1 targets Kar3 to its cytoplasmic binding sites [i.e., the MT plus ends] [Sproul et al. 2005] and, as shown in this study, the cytoplasmic side of the SPB through a Cik1-codependent interaction of its N-terminal domain with Spc72. Since Spc72 itself relocates to the half bridge during mating [Pereira et al. 1999], this leads to the reorganization of the entire cytoplasmic MT upon pheromone stimulation. At present, it is unclear how Spc72 binding to the SPB is regulated; one might speculate about a direct regulation of the Nucl1/Spc72 and/or Kar1/Spc72 interactions by kinases involved in pheromone signaling. Altogether, this sets the stage to transport the SPB—and with it, the attached nucleus—along MTs from the opposite SPB. This seems to be sufficient to promote nuclear congression in an asymmetric manner [MTs anchored at one SPB, while the other SPB is sliding along these MTs] [Fig. 5]. In wild-type cells, the process is symmetric, as both SPBs fulfill both roles simultaneously, increasing the overall robustness.

The realism of this model was proven, as nuclear congression occurred in silico with the very similar characteristics measured for the in vivo process. Nonetheless, functions that per se are not essential for congression may provide additional robustness to the process in vivo. First, the persistence of polarized actin cables could contribute to maintain the nucleus and the MTs oriented, such as the two SPBs remain facing each other until the MTs reach their opposite SPBs. Second, Kar3 is present also on the MT lattice, and its reported spindle function to regulate anti-parallel overlaps could promote such interactions also in karyogamy, as previously assumed [Rose 1996]. Similarly, Kar3 present at the MT plus ends could still promote direct interactions between plus ends, as suggested by the “plus end interaction model” [Molk et al. 2006]. The fact that we did not observe such MT–MT interactions in the limited numbers of tomograms that we generated does not mean that they never exist, but rather that they might be transient, occurring probably early in the process when these interactions may be promoting directional growth of MT toward opposite SPBs. The simulations indeed showed that direct Kar3-mediated MT–MT interactions could shorten the initial search phase and make the migration more robust by helping MTs find the opposite SPB.

Altogether, our data argue for a system of striking elegance in which only a few molecules are involved, but they are arranged in what seems to be the best possible configuration considering the task at hand. Moreover, the manner in which they promote direct transport of nuclei along MTs is similar to nuclear migration in higher cells.

In contrast to yeast, the animal centrosome is not passively embedded in the nuclear envelope but exists as
a dynamic organelle in the cytoplasm [Gönczy et al. 1999; Morris 2003]. In this context, the movement of nuclei often relies on the attachment of the nucleus to a polarized MT array through MT motor proteins. This is the case in mammalian epithelial cells in which kinesin 1 at the nuclear surface moves nuclei toward plus ends of polarized MTs to the base of the cell [Bacallao et al. 1989; Roux et al. 2009]. Similar mechanisms are involved in Xenopus female pronuclear movement, where the dynein-carrying female pronucleus tracks along MTs from the male-derived centrosome, thus pulling the two pronuclei together [Reinsch and Karsenti 1997]. The yeast nuclear congression process described here is strikingly similar. The nucleus is transported via the minus end-directed kinesin-14 motor Kar3 along an array of MT nucleated by the SPB of the opposite nucleus, independently of its own SPB nucleation properties. Interestingly, in yeast, the motors are only localized at the SPB and not all over the surface of the nucleus. The reason for this is likely to be a functional one: While in animal cells, bringing the nuclei together in any orientation is probably sufficient, in yeast, the correct orientation of the nuclei such that the two SPBs are able to meet and fuse seems to be a prerequisite. The different functional requirements are again reflected by the different molecular players: the SPB components in yeast and SUN–KASH bridges in animal cells [Starr and Fridolsson 2010].

Materials and methods

Strains and yeast methods

All yeast strains used in this study are derivatives of the S288c strains ESM356-1 [MATa] and ESM357-9 [MATa] [Knop and Schiebel 1998]. The genotypes of the yeast strains used in this work are listed in Supplemental Table S2. The plasmids used in this study are listed in Supplemental Table S3. Basic yeast methods and growth medium are described elsewhere (Sherman 1991). Chromosomal manipulations of yeast strains (C-terminal tagging, gene deletions, and promoter substitutions) were performed using PCR-amplified cassettes as described previously [Janke et al. 2004]. The N-terminal tagging of KAR3 was performed using seamless gene tagging by endonuclease-driven homologous recombination [Khmelinskii et al. 2011]. Quantitative mating essays were performed essentially as described [Pereira et al. 1999]. Two-hybrid interactions were determined using plasmids and strains described earlier [Geissler et al. 1996; Schramm et al. 2000].

Electron tomography

Cells of both mating types were grown in log phase in YPD medium (–0.5 × 10^7 cells per milliliter) and mixed in equal proportions. Mixed cells were collected onto a nitrocellulose membrane (0.45-μm pore size; Millipore) using vacuum filtration and placed on a YPD agar plate for 2 h at 30°C. Mating cells were cryoimmobilized by high-pressure freezing with a Leica EMPACT-2 [Leica Microsystems]. Samples were then freeze-substituted and plastic-embedded as described previously [Höög and Antony 2007] using 0.2% uranyl acetate, 0.1% glutaraldehyde, and 1% water in anhydrous acetone as a freeze substitution solution. Serial thick sections (300–320 nm thick) were cut using a Reichert Ultracut-E microtome (Leica Microsystems) and collected on Formvar-coated palladium–copper slot grids. Post-staining was performed with 4% uranyl acetate in 70% methanol for 10 min and subsequently with lead citrate (1.5% at pH 12) for 3 min. Tilted images were acquired at 300 kV from –60°C to +60°C with 1°C increments on a Tecnai F30 electron tomography microscope equipped with an Eagle 4K CCD camera (FEI: pixel size 1.499 nm at 15,500 magnification). Tomograms were then generated by R-weighted back projection, modeled, and analyzed using IMOD software as already described [Kremer et al. 1996].

Live-cell fluorescent microscopy

For imaging of mating yeast cells, cells of both mating types were grown in synthetic complete (SC)–complete medium to log phase (–0.5 × 10^7 cells per milliliter) and mixed together in equal proportions. Mixed cells were immobilized in glass-bottomed well chambers [LabTek 155411; Nalge Nunci International]. The chambers were pre-treated for at least 30 min with a 1:100 dilution of Bioconex in ethanol (UCT) followed by one ethanol and one water wash step, then incubated for >30 min with 5% Concanavalin A in water (C2010, Sigma) followed again by two washing steps with water. Following 1.5–2 h of incubation to allow mating to proceed, imaging was performed on a Deltavision RT wide-field microscope [Applied Precision] equipped with a uPlanApo 100× NA 1.35 oil immersion objective (Olympus), softWoRx software [Applied Precision], and a CoolSNAP HQ camera [Photometrics]. For α-factor treatment, MATα cells were grown in log phase in liquid SC medium and then immobilized in glass-bottomed well chambers, and α-factor was added to a final concentration of 10 μg/mL. Shmooing cells were imaged after 1.5–2 h of stimulation using the Deltavision RT microscope. Z-stacks (six planes at 0.5–0.6-μm interval, 1 × 1 binning) for each channel (GFP and mCherry) and a single plane bright-field image were acquired every 10 sec for 3–5 min. All movies show maximum projections of the Z-stacks at 10 frames per second (fps). For the purpose of visualization, images have been bleach-corrected by normalizing the intensities to the values at the beginning of the movie, and a Gaussian filter with a radius of 1 was applied.

FRAP

All imaging was performed at 30°C on a PerkinElmer Imagination Ultraview VoX spinning-disk confocal microscope equipped with a 100× 1.3 NA Plan immersion oil objective. GFP was excited using the 488-nm line of a 100-mW argon laser at 10% laser power [Spectra Physics]. A prebleach image was captured followed by bleaching of a circular area of 700 nm in diameter (four pulses of 1 msec of the 488-nm laser set to 60% intensity) followed by capturing of post-bleach images every 3 sec for 100 sec [exposure, 150 msec; binning, 2]. Six focal planes at 0.7-μm distance were captured using a Hamamatsu C9100-50 camera. The centroid of the GFP-Kar3 spot has been tracked using the tracking algorithm from Shalzarini and Koumoutsakos [2005]. Fitting of a Gaussian to the fluorescent signal yielded a standard deviation σ of ~1.5 pixels. The region of interest was taken in the 2σ limit. We thus computed the intensity of an area of 6 × 6 pixels around the centroids of the bleached SPBs. This has been bleach-corrected using the total GFP-Kar3 fluorescence of other cells in the field of view as a reference.

Protein localization at the SPB

Cells were imaged on a Zeiss LSM 780 using a 63×1.4 NA PlanApo immersion oil objective at 30°C. GFP and mCherry
were excited simultaneously with a 488-nm argon laser and a 561-nm DPSS laser, respectively. The spectral array detector was used to image the green and red channels simultaneously. We acquired 26 planes with 0.2-μm distance. The position of the proteins at the SPB was determined by finding the peak of a Gaussian fitted to the fluorescence signal in the plane with maximal intensity. The distal position of each nucleus with respect to Spc42-mCherry served as a reference coordinate for the nucleus.

Quantification of GFP-Kar3 at the SPB and measurement of nuclear congression kinetics

Images were acquired using a wide-field Olympus IX81 with a 100×1.4 NA PlanApo oil objective and a Hamamatsu OrcaER camera (binning of 2, pixel size, 0.13 μm). For the quantification of the GFP signal of GFP-Kar3, MATα and MATa GFP-KAR3-expressing cells were mixed and allowed to mate as described before. After 1.5 h, MATα NU2F2-GFP PIL1-mCherry-expressing cells were added to the glass-bottomed well chambers and allowed to settle for 10 min. The Pil1-mCherry signal was used to identify these cells. Per field of view, 31 planes at a 0.2-μm distance (200 msec exposure for GFP, 100 msec for mCherry) were acquired. Quantification was done as described by Joglekar et al. (2006). The number of GFP molecules present in the GFP-Kar3 spot was calculated by comparison with the Nu2-F2-GFP spot in metaphase cells, which contains on average 385.6 molecules of GFP (Lawrimore et al. 2011).

To determine the SPBs’ nuclear congression dynamics, cells of both mating types expressing SPc42-GFP (or CNM67-GFP-KAR3) and mCherry-TUB1 were used. To determine the time point of cell fusion, one mating partner additionally expressed soluble cytoplasmic TagBFP from the GPD promoter. Congresion kinetics were acquired. Quantification was done as described above. Tracking of the position of the SPBs was conducted as described above.

First principles modeling of MT and nuclear dynamics

In silico nuclear congression simulations were carried out using Cytosim, a program developed for stochastic simulations of cytoskeletal fibers, diffusible activities, and flexible objects in a confined environment. The simulation is based on Brownian dynamics, and the system properties emerge from real amounts of constituents following a stochastic description of the most relevant physical processes [Nedelec and Foethke 2007]. The nucleus is spherical, but the SPB is another smaller sphere attached on its side with three Hookean springs. This setup represents the nuclear protrusions observed in the tomograms. The MTs are anchored at the SPB and have a restricted rotational freedom at their SPB attachment point so that, on average, they grow away from the nucleus from which they were nucleated. MTs are modeled as dynamic growing fibers that stochastically switch to shrinkage. Below a minimal length, MTs are rescued and switch back to growth. In the model, we restrict ourselves to the situation just after cell fusion and assume that nuclei and MTs are properly aligned toward the shmoo as observed [Maddox et al. 2003]. The initial distance of SPB to neck was 3.08 ± 0.61 μm for wild type and 2.4 ± 0.54 μm for CNM67-GFP-KAR3 kar3Δ to match the measured distances at cell fusion. MTs were straight in the growing or shrinking states with equal probability, and their lengths were randomly distributed with the constraint that they should fit in the cell before fusion. The model parameters for the dynamics of MTs, the Kar3 minus end-directed motors, and the geometry of the mating cells have been obtained from the literature, the tomograms presented here, and the dynamics of GFP-Kar3 during congression [Supplemental Table S1]. To simulate the MT signal in light microscopy, we approximated the point spread function (PSF) of the microscope with a Gaussian of width

\[ \sigma_{xy} = \sqrt{\left(\frac{0.25a}{NA}\right)^2 + \frac{1}{12a^2}}, \]

where NA = 1.4 is the numerical aperture, \( a = 0.13 \) μm is the pixel size, and \( \lambda \) is the emission wavelength [Mortensen et al. 2010; Stallinga and Rieger 2010]. Model fluorophores, uniformly distributed on MTs, are convolved with this function.

Acknowledgments

We thank the members of the Antony, Knop, and Nédélec groups and the EMBL Electron Microscopy Core Facility for helpful discussions and technical support. We acknowledge the people in the Nédélec group involved in the development of Cytosim. Kornelius Schweinfurth is acknowledged for help in tomography modeling. We thank Susanne Trautmann, Celine Maeder, and Matthias Meurer for plasmids. We acknowledge Tomo Tanaka for discussions. We are grateful to Elmar Schiebel for plasmids, discussion, and comments on the manuscript. We also thank Charlotte Funaya for critical reading. A.Z.P. is supported by a post-doctoral grant from BIOMS (Center for Modelling and Simulation in the Biosciences in Heidelberg). C.A., M.K. and F.N. designed the project. R.G. and A.Z.P. together did the light microscopy experiments and image analysis. R.G. performed yeast genetic experiments and electron microscopy, and A.Z.P. performed the first principles modeling. M.K. and R.G., helped by A.Z.P., wrote the manuscript with input from C.A. and F.N. All authors discussed the results and commented on the manuscript.

References


Elliott S, Knop M, Schlenstedt G, Schiebel E. 1999. Spc29p is a component of the Spc110p subcomplex and is essential for

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Sharma C, Elliott S, Shevchenko A, Schiebel E. 2000. The Bbp1p–Mps2p complex connects the SPB to the nuclear
envelope and is essential for SPB duplication. EMBO J 19:421–433.