

# Structural Mutants of the Spindle Pole Body Cause Distinct Alteration of Cytoplasmic Microtubules and Nuclear Dynamics in Multinucleated Hyphae

Claudia Lang,\* Sandrine Grava,\* Mark Finlayson,\* Rhonda Trimble,<sup>†</sup>  
Peter Philippsen,\* and Sue L. Jaspersen<sup>†‡</sup>

\*Department of Molecular Microbiology, Biozentrum University of Basel, 4056 Basel, Switzerland; <sup>†</sup>Stowers Institute for Medical Research, Kansas City, MO 64110; and <sup>‡</sup>Department of Molecular and Integrative Physiology, University of Kansas Medical Center, Kansas City, KS 66160

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In the multinucleate fungus *Ashbya gossypii*, cytoplasmic microtubules (cMTs) emerge from the spindle pole body outer plaque (OP) in perpendicular and tangential directions. To elucidate the role of cMTs in forward/backward movements (oscillations) and bypassing of nuclei, we constructed mutants potentially affecting cMT nucleation or stability. Hyphae lacking the OP components AgSpc72, AgNud1, AgCnm67, or the microtubule-stabilizing factor AgStu2 grew like wild-type but showed substantial alterations in the number, length, and/or nucleation sites of cMTs. These mutants differently influenced nuclear oscillation and bypassing. In *Agspc72Δ*, only long cMTs were observed, which emanate tangentially from reduced OPs; nuclei mainly moved with the cytoplasmic stream but some performed rapid bypassing. *Agnud1Δ* and *Agcnm67Δ* lack OPs; short and long cMTs emerged from the spindle pole body bridge/half-bridge structures, explaining nuclear oscillation and bypassing in these mutants. In *Agstu2Δ* only very short cMTs emanated from structurally intact OPs; all nuclei moved with the cytoplasmic stream. Therefore, long tangential cMTs promote nuclear bypassing and short cMTs are important for nuclear oscillation. Our electron microscopy ultrastructural analysis also indicated that assembly of the OP occurs in a stepwise manner, starting with AgCnm67, followed by AgNud1 and lastly AgSpc72.

## INTRODUCTION

Microtubule-organizing centers (MTOCs) are a structurally diverse class of organelles involved in cell division, intracellular trafficking, cytoplasmic organization, and motility. Defined by the localization of  $\gamma$ -tubulin and its associated proteins, the function of MTOCs in nucleation and anchorage of microtubules is highly conserved in all eukaryotes. Centrosomes are the primary MTOC in metazoans, whereas nuclear-associated spindle pole bodies (SPBs) function as MTOCs in fungi. In the budding yeast *Saccharomyces cerevisiae*, the SPB serves as the sole site of microtubule nucleation, forming both nuclear microtubules involved in spindle assembly and chromosome segregation and cytoplasmic microtubules (cMTs) involved in nuclear migration and spindle positioning (reviewed in Jaspersen and Winey, 2004). The multinucleate filamentous fungus *Ashbya gossypii* also contains nuclear-associated SPBs as its only MTOC (Lang *et al.*, 2010).

Structural analysis of the multilayered *A. gossypii* SPB from wild-type cells by using electron microscopy (EM) revealed considerable similarity to the SPB of *S. cerevisiae* (Lang *et al.*, 2010). However, structural differences with functional implications were found at the cytoplasmic side of the *A. gossypii* SPB. Two types of cMTs emerge from a spherical outer plaque (OP), one type in perpendicular and the other type in tangential orientations. The perpendicular cMTs are generally short and make connections with the cortex, similar to budding yeast cMTs in G1 cells (Carminati and Stearns, 1997; Shaw *et al.*, 1997). The tangential class of cMTs is not found in budding yeast. These very long cMTs bypass other nuclei in the multinucleated hyphae. A model for the role of both types of cMTs in nuclear movements was proposed in Lang *et al.*, 2010 based on EM and live cell imaging experiments. Long tangential cMTs were predicted to be important for long-range nuclear bypassing events and the shorter cortex-connected cMTs were predicted to play a role in short-range nuclear oscillations. To test this model, it is necessary to find and investigate mutants with altered cMT nucleation and nuclear migration behavior. Based on the evolutionary relationship between *A. gossypii* and *S. cerevisiae* (Dietrich *et al.*, 2004), it is highly likely that both organisms use similar proteins to nucleate and anchor microtubules. Therefore, genes deleted for analysis of cMT function in multinucleated *A. gossypii* hyphae were selected based on extensive knowledge of *S. cerevisiae* SPBs.

In budding yeast, three SPB substructures are directly involved in binding of the  $\gamma$ -tubulin complex and thus mi-

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Address correspondence to: Sue L. Jaspersen (slj@stowers.org).

Abbreviations used: cMT, cytoplasmic microtubule; EM, electron microscopy; IL, intermediate layer; IP, inner plaque; MEN, mitotic exit network; MTOC, microtubule-organizing center; OP, outer plaque; SPB, spindle pole body.

**Table 1.** Amino acid sequence comparison of SPB components

<i>S. cerevisiae</i> SPB component <sup>a</sup>	<i>S. cerevisiae</i> SPB localization	Role in SPB function in <i>S. cerevisiae</i>	<i>A. gossypii</i> orthologue	Protein length in amino acids ( <i>S.c./A.g.</i> ) <sup>b</sup>	% identity <sup>b</sup>
Tub4	$\gamma$ -Tubulin complex	Microtubule nucleation	AgTub4	473/470	56
Spc97	$\gamma$ -Tubulin complex	Microtubule nucleation	AgSpc97	823/836	30
Spc98	$\gamma$ -Tubulin complex	Microtubule nucleation	AgSpc98	846/845	41
Spc72	OP, HB	$\gamma$ -tubulin complex binding protein	AgSpc72	622/795	22
Nud1	OP	MEN signaling	AgNud1	851/757	26
Cnm67	IL1, OP	Spacer, anchors OP to CP	AgCnm67	581/862	18
Spc42	CP, IL2	Structural SPB core	AgSpc42	363/314	28
Spc29	CP	Structural SPB core	AgSpc29	253/293	20
Cmd1	CP	Spc110 binding protein	AgCmd1	147/147	95
Spc110	CP to IP	$\gamma$ -Tubulin complex binding protein	AgSpc110	944/852	26
Mps2	SPB periphery	SPB insertion	AgMps2	387/338	19
Ndc1	SPB periphery	SPB insertion	AgNdc1	655/591	36
Bbp1	SPB periphery	SPB core to HB membrane linker	AgBbp1	385/353	26
Nbp1	SPB periphery	SPB core to HB membrane linker	AgNbp1	319/328	28
Cdc31	HB	SPB duplication	AgCdc31	161/172	71
Sfi1	HB	SPB duplication	AgSfi1	946/991	24
Kar1	HB	SPB duplication	AgKar1	433/330	23
Mps3	HB	SPB duplication	AgMps3	682/616	35

<sup>a</sup> SPB components are defined as proteins required to maintain the structural integrity of the organelle (Jaspersen and Winey, 2004).

<sup>b</sup> Based on gene information in the *S. cerevisiae* and *A. gossypii* databases. Percentage of identity was determined along the entire length as described in *Materials and Methods*.

microtubule nucleation. The  $\gamma$ -tubulin complex is tethered to the nuclear face of the SPB, known as the inner plaque (IP), by Spc110 and nucleates the intranuclear microtubules that form the spindle (Kilmartin and Goh, 1996; Spang *et al.*, 1996; Knop and Schiebel, 1997; Nguyen *et al.*, 1998; Pereira *et al.*, 1998). A different  $\gamma$ -tubulin complex binding protein, Spc72, is used for tethering Tub4, Spc97, and Spc98 to the cytoplasmic side of the SPB. The site of Spc72 binding and cMT nucleation changes throughout the cell division. During most of the cell cycle, Spc72 is present at the SPB OP but during G1 phase it can be associated with the half-bridge or bridge region (Rout and Kilmartin, 1990; Knop *et al.*, 1997; Knop and Schiebel, 1998). Other components required for formation of the OP and anchoring of cMTs include the intermediate layer (IL)1 to OP spacer Cnm67 and the OP protein Nud1 (Brachat *et al.*, 1998; Elliott *et al.*, 1999; Gruneberg *et al.*, 2000; Schaerer *et al.*, 2001; Muller *et al.*, 2005). In contrast to Spc72, which can be found associated with the half-bridge or OP depending on the cell cycle stage, Cnm67 and Nud1 are localized exclusively to the OP of the SPB (Brachat *et al.*, 1998; Adams and Kilmartin, 1999; Pereira *et al.*, 1999).

Although most SPB components are essential for viability in budding yeast due to their role in SPB duplication and bipolar spindle formation, deletion analysis showed that components of the outer layers of the SPB are not absolutely required for viability in all strain backgrounds (Brachat *et al.*, 1998; Soues and Adams, 1998; Hoepfner *et al.*, 2000, 2002). However, *cnm67* $\Delta$  or *spc72* $\Delta$  cells often lack a nucleus or carry two or more nuclei due to defects in nuclear migration and spindle positioning, processes that both require cMTs. Survival of *cnm67* $\Delta$  cells, which lack an OP and the cMTs normally formed at this structure, results from a rescue pathway involving microtubule nucleation from the half-bridge (Brachat *et al.*, 1998; Hoepfner *et al.*, 2000). Certain

temperature-sensitive mutants such as *nud1-2* share a similar phenotype to *cnm67* $\Delta$  cells (Gruneberg *et al.*, 2000), suggesting that it is also essential for OP formation and microtubule nucleation. Analysis of *NUD1* is complicated by the fact that this SPB component serves as a scaffold for a signaling pathway that monitors spindle positioning and controls mitotic exit, known as the mitotic exit network (MEN; reviewed in Hoyt, 2000; Pereira and Schiebel, 2001; Stegmeier and Amon, 2004). Segregation of chromosomes to the daughter cell is critical in budding yeast, so deletion of *NUD1* or most MEN components results in lethality. The SPB may also be a loading or storage site of Stu2, an essential, conserved microtubule-plus end binding protein of the XMAP215/Dis1 family that regulates both nuclear and cMT dynamics (Kosco *et al.*, 2001; Severin *et al.*, 2001; Pearson *et al.*, 2003; van Breugel *et al.*, 2003; Usui *et al.*, 2003; Al-Bassam *et al.*, 2006).

*A. gossypii* carries syntenic homologues for these SPB genes (Table 1). However, some of the encoded orthologues have <20% sequence identity, and it is unclear whether and how differences in the primary sequence translate into changes in SPB structure and cMT nucleation or anchorage needed to coordinate movements of nuclei in a multinucleated cytoplasm. To better understand which type of cMT controls nuclear oscillation or nuclear bypassing in *A. gossypii*, we constructed and analyzed deletions of *A. gossypii* genes orthologous to SPB genes of budding yeast. We found that some deletions had an analogous phenotype in *A. gossypii* as they did in *S. cerevisiae*, whereas several other deletions resulted in unexpected and novel phenotypes, which allowed us to assign functions to the two types of cMTs and to present the ultrastructure of SPBs and attached cMTs in *Agspc72* $\Delta$ , *Agstu2* $\Delta$ , *Agcnm67* $\Delta$  and *Agnud1* $\Delta$  mutants.

## MATERIALS AND METHODS

### *A. gossypii* Media and Growth Conditions

*A. gossypii* media and culturing are described in Ayad-Durieux *et al.* (2000) and Wendland *et al.* (2000), and strains are listed in Supplemental Table S1.

### Plasmid and Strain Construction

Plasmids generated and used in this study are described below. All DNA manipulations were carried out according to Sambrook and Russell (2001) with *Escherichia coli* DH5 $\alpha$ F' as host (Hanahan, 1983). Polymerase chain reaction (PCR) amplification was performed using standard methods with *Taq* DNA polymerase, Expand High Fidelity PCR system, or the Expand Long Template PCR system (Roche Diagnostics, Indianapolis, IN). Oligonucleotides are listed in Supplemental Table S2 and were synthesized by Microsynth (Balgach, Switzerland).

*A. gossypii* deletion mutants were made using the PCR-based one-step gene targeting approach with heterologous selection markers (Wendland *et al.*, 2000). The deletion cassettes were amplified from a plasmid containing the *GEN3* (Wendland *et al.*, 2000) cassette that mediates resistance to G418, using 'gene name'\_NS1/F2 oligonucleotide pairs. Correct integration of the deletion cassettes was verified with oligonucleotide primer pairs 'gene name'\_A1/Gen2\_A2 (N terminus) and 'gene name'\_A4/Gen2\_A3 (C terminus). Three independent transformants were characterized for each mutant. Transformation of multi-nucleate mycelium leads to heterokaryotic cells, which contain a mixture of transformed and wild-type nuclei. For subsequent analysis, homokaryotic mycelia were obtained by isolating and growing single spores. To evaluate phenotypes of lethal mutants or mutants with sporulation deficiency, spores from heterokaryotic mycelium were germinated and analyzed under selective conditions (200  $\mu$ g/ml G418; Sigma-Aldrich, St. Louis, MO).

### Fluorescence Microscopy and Image Processing

DNA (Hoechst) and immunofluorescence stainings were performed as described previously (Ayad-Durieux *et al.*, 2000; Gladfelter *et al.*, 2006). Rat anti- $\alpha$ -tubulin (YOL1/34; Serotec, Oxford, United Kingdom) was used at a 1:25 dilution and Alexa Fluor 568 goat anti-rat immunoglobulin (IgG; Invitrogen, Carlsbad, CA) at a 1:200 dilution.

An Axioplan2 microscope equipped with the objectives Plan-Apochromat 100 $\times$ /1.40 numerical aperture (NA) Oil differential interference contrast (DIC) and Plan-Apochromat 63 $\times$ /1.40 NA Oil DIC (Carl Zeiss, Feldbach, Switzerland) and appropriate filters (Carl Zeiss and Chroma Technology, Brattleboro, VT) was used for microscopy. The light source for fluorescence microscopy was either a 75-W XBO lamp (Osram, Augsburg, Germany), controlled by a MAC2000 shutter and filter wheel system (Ludl Electronics, Hawthorne, NY) or a Polychrome V monochromator (TILL Photonics, Gräfelfing, Germany). Images were acquired at room temperature using a cooled charge-coupled device camera (CoolSNAP HQ; Photometrics, Tucson, AZ) with MetaMorph 6.2r5 software (Molecular Devices, Sunnyvale, CA). Out-of-focus shading references were used for DIC image acquisitions.

For time-lapse image acquisition, a glass slide was covered with 1 ml of *A. gossypii* minimal medium containing 1% agarose. Once the medium had solidified, either small pieces of mature mycelium from the border of 3-d-old *A. gossypii* colonies or young mycelia cultured in liquid medium were spotted onto the slides. Seventy microliters of liquid minimal medium was added to the mycelia before cells were covered with a coverslip and incubated for at least 1 h before image acquisition. For still images, multiple planes with a distance between 0.3 and 1  $\mu$ m in the z-axis were taken.

Image processing was performed with MetaMorph 6.2r5 software. Z-stacks were deconvolved with Nearest Neighbor and compressed by maximum or average projection with Stack Arithmetic. Brightness and contrast were adjusted using Scale Image. Images were colored and overlaid by using Overlay Images and exported from MetaMorph as 8-bit grayscale or RGB TIFF files. Z-Stacks and time-lapse picture series were converted to QuickTime H.264 movies with Quicktime Player Pro (Apple Computer, Cupertino, CA).

### Transmission Electron Microscopy

Spores were grown for 10–14 h in liquid AFM to give rise to small mycelia containing no >100 nuclei. Samples were frozen on the Leica EM-Pact at ~2050 bar and then transferred under liquid nitrogen into 2% osmium tetroxide/0.1% uranyl acetate/acetone and transferred to an automated freeze-substitution apparatus (AFS) (Leica, Wetzlar, Germany). The freeze substitution protocol was as follows: -90°C for 16 h, up 4°C/h for 7 h, -60°C for 19 h, up 4°C/h for 10 h, and -20°C for 20 h. Samples were removed from the AFS and placed in the refrigerator for 4 h and then allowed to incubate at room temperature for 1 h. Samples went through three changes of acetone over 1 h and were removed from the planchettes. Then, they were embedded in acetone/Epon mixtures to final 100% Epon over several days in a stepwise procedure as described previously (McDonald, 1999). We cut 60-nm serial thin sections on a UC6 ultramicrotome (Leica), stained them with uranyl acetate and Sato's lead, and imaged them on a Spirit transmission electron microscope (FEI Technai, Hillsboro, OR). Serial section images were aligned using AutoAligner (Bitplane, Zurich, Switzerland). For the *AgNUD1* deletion

and some samples of the *AgCNM67* deletion, mycelium of the border of 3-d-old colonies was frozen and subsequently treated as described above. We could not detect any differences in the SPB structure between either method of sample preparation.

### Bioinformatic Analysis

Nuclear localization signal (NLS) search was performed with PredictNLS (<http://cubic.bioc.columbia.edu/predictNLS/>) and Prosite (<http://www.expasy.org/prosite/PS50079>). Protein alignments were performed with sequences retrieved from the *Ashbya* Genome Database (<http://agd.vital-it.ch/>; Gattiker *et al.*, 2007) and the *Saccharomyces* Genome Database (<http://www.yeastgenome.org/>) by using the EMBOSS Pairwise Alignment Algorithms (Blosum62 Matrix, gap open 10, gap extend 0.5).

## RESULTS

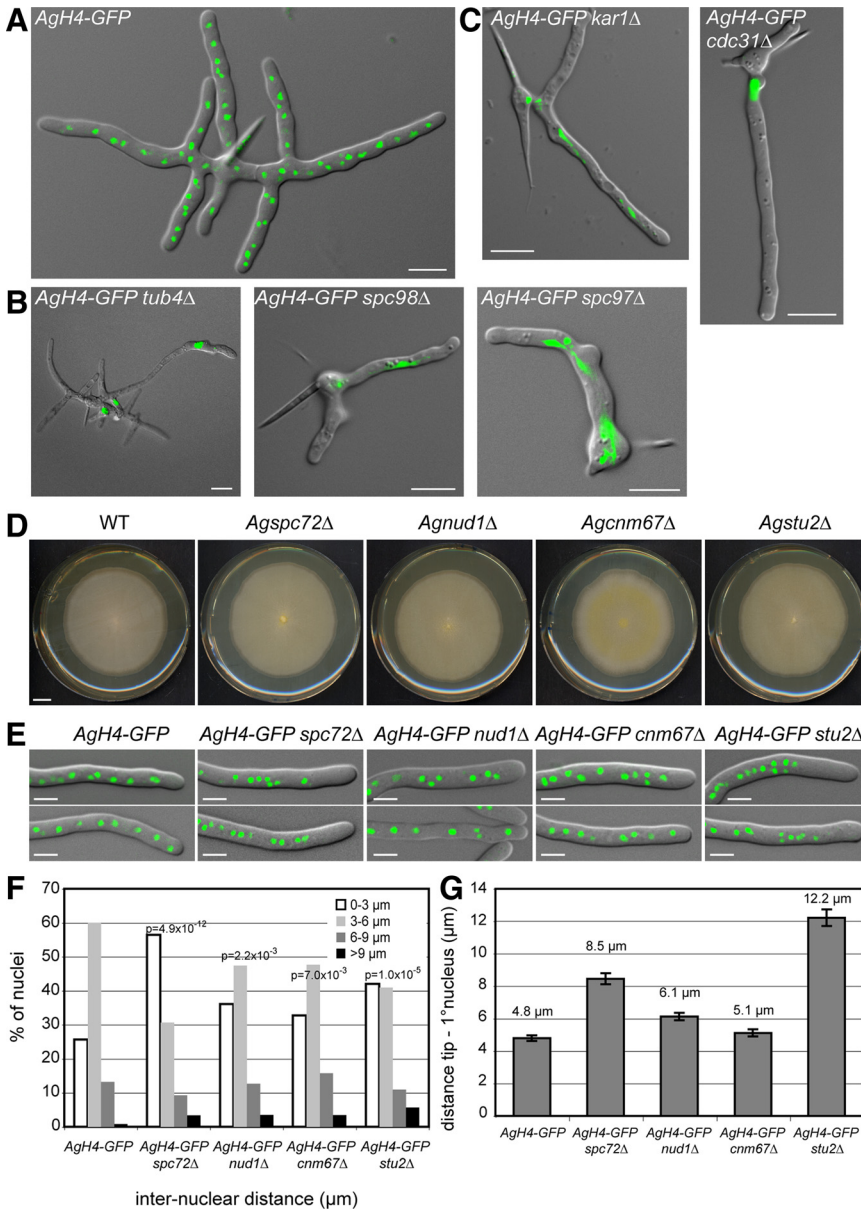
### Essential and Nonessential SPB Components in *A. gossypii*

We searched the *A. gossypii* genome for homologues of genes that encode components of the evolutionary related budding yeast SPB. Sequence analysis revealed that the *A. gossypii* genome encodes syntenic homologues for all 18 mitotic *S. cerevisiae* SPB components (Table 1). Most orthologous proteins share only 20–40% identity. Notable exceptions are Tub4 ( $\gamma$ -tubulin) and the calcium-binding proteins Cmd1 (calmodulin) and Cdc31 (centrin), which are 56, 95, and 71% identical, respectively. These proteins are conserved components of many MTOCs, including SPBs and centrosomes (Jaspersen and Winey, 2004).

Given the structural similarities between the *A. gossypii* and *S. cerevisiae* SPB (Lang *et al.*, 2010) but the relatively low level of sequence conservation of its constituent parts, we were interested to test functional conservations using a gene deletion approach. We asked whether elimination of core cytoplasmic components had similar effects on SPB structure, microtubule nucleation, and nuclear migration in multinucleated hyphae as in budding yeast cells or whether some deletions caused unexpected phenotypes due to the evolutionary adaptation of mechanisms of nuclear migration in both systems. A priori, it cannot be excluded, that an orthologue is essential in *S. cerevisiae* but nonessential in *A. gossypii*, so deletion analysis of SPB components in *A. gossypii* could result in structural and functional insight of mutant SPBs not possible in budding yeast. Specifically, we created deletion mutants in components of the  $\gamma$ -tubulin complex, the half-bridge, and the OP because these SPB substructures have well-documented roles in cMT organization as well as nuclear migration and positioning in *S. cerevisiae* (Rose and Fink, 1987; Geissler *et al.*, 1996; Spang *et al.*, 1996; Knop *et al.*, 1997; Brachat *et al.*, 1998; Chen *et al.*, 1998; Pereira *et al.*, 1999; Gruneberg *et al.*, 2000; Hoepfner *et al.*, 2000, 2002; Usui *et al.*, 2003). In our analysis of *A. gossypii* SPB deletion mutants, we mainly focused on nuclear migration dynamics, formation of cMTs, and the structure of the mutant SPBs with the aim of providing a mechanistic model for long-range nuclear migration within multinucleate hyphae.

We found that *A. gossypii* genes encoding components of the  $\gamma$ -tubulin complex (*AgTUB4*, *AgSPC97*, and *AgSPC98*) are all essential. Visualization of nuclei in germinated spores lacking *AgTUB4*, *AgSPC97*, or *AgSPC98* by using histone H4-green fluorescent protein (*AgH4-GFP*) revealed a clear nuclear division defect: nuclear density was decreased ~10-fold compared with wild-type ( $n > 100$ ) and nuclei were elongated or fragmented (Figure 1, A and B). The  $\gamma$ -tubulin complex probably nucleates microtubules at both the IP and OP of the SPB. Because cMTs are dispensable for hyphal growth of *A. gossypii* (Lang *et al.*, 2010), the lethality of these mutants is probably due to defects in spindle





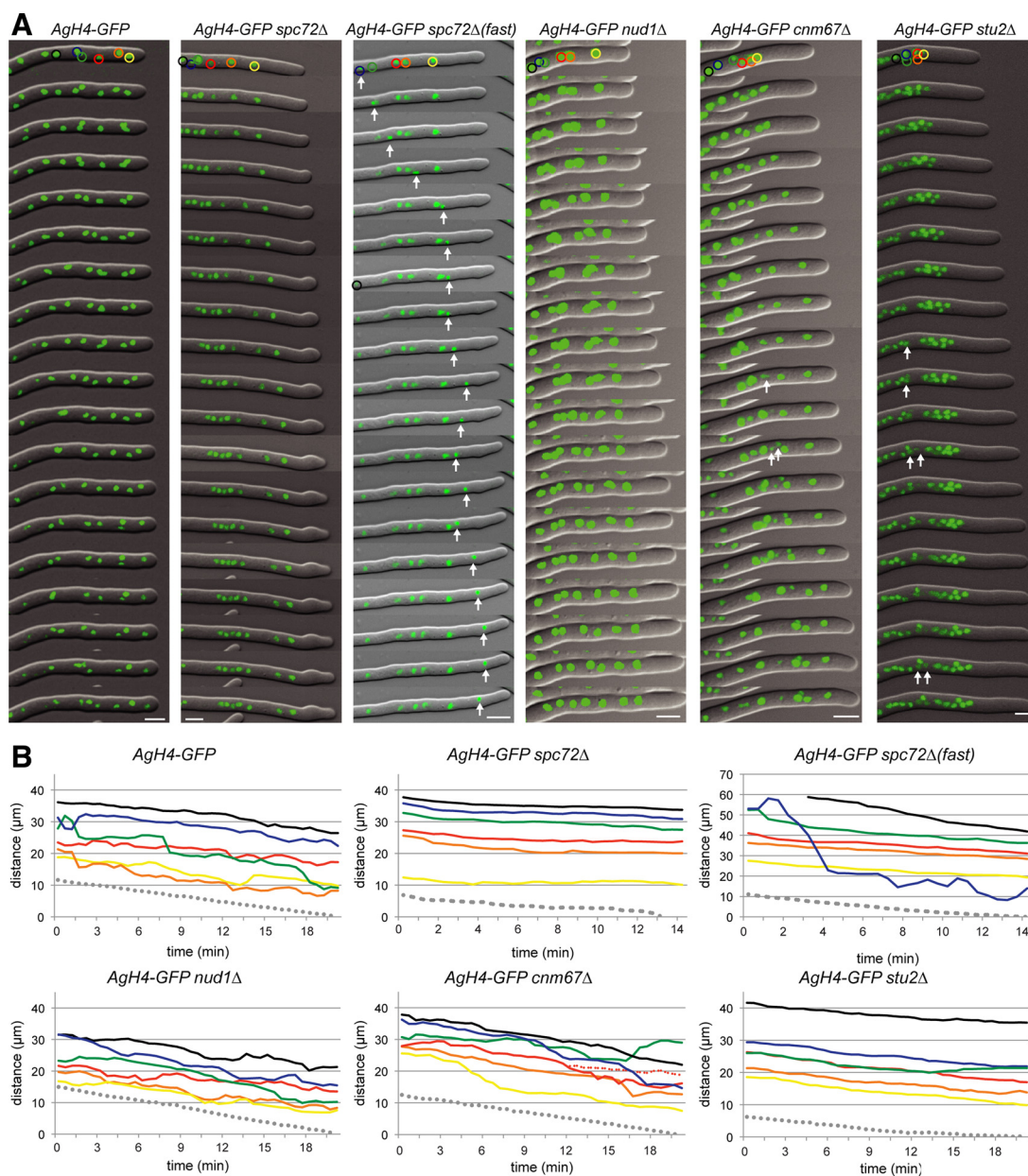
**Figure 1.** Deletion of *A. gossypii* SPB components. Overlays of DIC image and AgH4-GFP signals from cells grown for ~12 h at 30°C. (A) A typical wild-type mycelium at this stage contains multiple branches and 50–100 nuclei. (B) Terminal phenotype of mutants lacking components of the  $\gamma$ -tubulin complex. (C) Terminal phenotype of mutants lacking half-bridge components. Some mutants shown in B and C arrest as small mycelium with up to 12 branches containing a few nuclei and did not arrest as germlings with one nucleus or two nuclei. This is due to a maternal effect, where remnants of the wild-type protein are packed into mutant spores (Gladfelter *et al.*, 2006). In *A. gossypii*, targeted deletions are performed with young mycelium containing ~20 haploid nuclei that initially generate heterokaryotic strains: mixtures of nuclei carrying either the wild type or the deletion allele. On starvation, haploid mononucleate spores are produced in the heterokaryotic hyphae, and a substantial part of the mutant spores contain wild-type (maternal) protein, which can account for early growth and nuclear division in otherwise fatal deletions. Bars, 10  $\mu$ m. (D) Radial growth of wild-type and deletion mutants on solid medium during 7 d of incubation at 30°C. Bar, 1 cm. (E) Overlays of DIC image and AgH4-GFP signals from hyphae showing nuclear distributions in wild-type and mutant strains. Bars, 5  $\mu$ m. (F) Distribution of distances between adjacent nuclei was plotted using distance measurements between the nuclei in the first 50  $\mu$ m of at least 20 different hyphae for each strain ( $n > 200$  for each strain). The center of the GFP signal was used as the central point of the nucleus. Confidence values ( $p$ ) for the  $\chi^2$  test were calculated for each data set between wild-type and mutant distributions. (G) Distance between the hyphal tip and the first nucleus was determined. Error bars represent the SE of the mean,  $n > 60$  for each strain.

microtubule nucleation at the IP rather than nucleation of cMTs at the OP.

During G1 phase of the *S. cerevisiae* cell cycle, cMTs are nucleated on the half-bridge in a Kar1-dependent manner (Byers and Goetsch, 1975; Pereira *et al.*, 1999). The ability to form microtubules at the half-bridge is essential for survival of SPB OP mutants such as *cnm67Δ* (Brachat *et al.*, 1998; Hoepfner *et al.*, 2000). In *A. gossypii*, hyphae lacking the half-bridge component AgKar1 stopped growing as small mycelium containing up to 10 nuclei (Figure 1C). Thus, AgKAR1 is an essential gene. Dependent on the remnant AgKar1 proteins in mutant spores, SPBs can still duplicate two to three times, permitting a few nuclear divisions. The fact that deletion of *AgCDC31*, encoding another half-bridge component presumably not involved in cMT organization, resulted in the same terminal phenotype (cessation of growth as small mycelium; Figure 1C), suggests that lethality in cells lacking AgKAR1 is due to a general SPB duplication defect rather than a specific effect on cMTs.

Deletions of *AgSPC72*, *AgNUD1*, *AgCNM67*, and *AgSTU2* did not affect radial growth of colonies, a sensitive method to test for reduced hyphal growth (Figure 1D). *S. cerevisiae* orthologues of the first three genes encode OP components, the orthologue of *AgSTU2* encodes an OP-associated MT-stabilizing protein, and deletions of these genes are either lethal or cause reduced growth in *S. cerevisiae* (Brachat *et al.*, 1998; Chen *et al.*, 1998; Knop and Schiebel, 1998; Soues and Adams, 1998; Adams and Kilmartin, 1999; Hoepfner *et al.*, 2000, 2002; Usui *et al.*, 2003; Al-Bassam *et al.*, 2006). Even though polar growth was not affected in the four *A. gossypii* deletions, alterations in nuclear distributions were noted in each (Figure 1, E–G).

The deletions are expected to alter the formation of microtubules at the cytoplasmic side of SPBs and microtubule stability thus affecting nuclear mobility. We therefore monitored the migration of H4-GFP labeled nuclei in wild-type and the four deletions, quantified the frequencies of forward and backward movements (oscillations) and bypassing of nuclei, visualized MTs *in vitro* by immunofluorescence micros-



**Figure 2.** Time-lapse analysis of nuclear migration in different mutants. (A) Overlays of DIC and AgH4-GFP signals from time-lapse video imaging of wild-type (Supplemental Movie S1), *Agspc72Δ* (Supplemental Movie S2), *Agnud1Δ* (Supplemental Movie S4), *Agcnm67Δ* (Supplemental Movie S5), and *Agstu2Δ* (Supplemental Movie S6) hyphae. Images were captured every 30 s, and 1-min interval frames are shown. Migration of the first six nuclei were tracked and are shown in the schematic. In *Agspc72Δ(fast)* (Supplemental Movie S3), images from each 30-s time point are shown for the first 7 min followed by 1-min interval frames; the rapidly moving nucleus is indicated by an arrow. Nuclei undergoing mitosis are indicated by arrows in *Agcnm67Δ* and *Agstu2Δ*. Bars, 5  $\mu\text{m}$ . (B) Positions of the first six nuclei in each hyphae and the hyphal tip (gray dotted line) were tracked throughout each time course and are plotted. In wild-type, the nuclei were observed to undergo bypassing and oscillations. In *Agspc72Δ(fast)*, one nucleus (blue) moves rapidly toward the tip, thereby bypassing four other nuclei in <5 min and traveling distances up to 8.9  $\mu\text{m}$  within a 30-s interval (1.5–2 min). The other nuclei in this hypha as well as in the other *Agspc72Δ* and *Agstu2Δ* mutants move toward the tip with the cytoplasmic stream without undergoing any bypassing or oscillation.

copy and in vivo using GFP-AgTub1 and finally used EM to determine the structure of SPBs in the mutants and their ability to nucleate perpendicular and tangential cMTs. Because of the complexity of this analysis we have documented the nuclear mobilities for all mutants in one figure and one table to allow for direct comparisons (Figure 2; Table 2), and we compiled separately for each mutant images of cMTs and SPBs (Figures 3–6). In the following paragraphs, we present and discuss the data for each of the four mutants individually, starting each

time with a functional description of the *S. cerevisiae* orthologue. The structural and functional results of our studies will be summarized at the end (Figures 7–9).

#### *Agspc72Δ* Lacks Nuclear Oscillations but Occasionally Shows Bypassing

Spc72 serves as the cytoplasmic  $\gamma$ -tubulin complex anchor in *S. cerevisiae* (Knop and Schiebel, 1998; Pereira *et al.*, 1999) and a deletion of the *ScSPC72* gene is lethal or yields in some



**Table 2.** Nuclear oscillations and bypassing in OP and *Agstu2Δ* mutants

Strain <sup>a</sup>	Distance migrated ( $\mu\text{m}$ )/30 s				Nuclear oscillations <sup>b</sup>		Nuclear bypassing		
	Avg.	max	n	p <sup>c</sup> value	Forward movement	Backward movement	Events	Total nuclei	%
Wild-type	0.51	2.82	1264		101	41	15	48	31
<i>spc72Δ</i>	0.35 (2.12)	1.39 (8.91)	1177 (82) <sup>d</sup>	<0.001 <0.001	48	0	4 (8) <sup>e</sup>	42	10
<i>nud1Δ</i>	0.45	2.3	861	<0.01	81	28	8	36	22
<i>cnm67Δ</i>	0.56	3.11	770	<0.05	138	28	17	25	68
<i>stu2Δ</i>	0.36	1.23	240	<0.001	9	0	0	32	0

<sup>a</sup> Background AgH4-GFP.

<sup>b</sup> The movements of the five most apical nuclei in four different hyphae (1 hypha for *stu2Δ*) were followed >40 or >30-s time intervals. Forward or backward displacements  $\geq 0.75 \mu\text{m}$  (approximately half the diameter of a nucleus) within 30 s were scored.

<sup>c</sup> Statistical significance between mutant and wild-type values was calculated using an unpaired Student's *t* test, and the resulting p value is shown. We also compared different mutant data sets with each other and all average values are highly statistically significant except for the slight difference in the average values from *Agspc72Δ* and *Agstu2Δ* mutants ( $p > 0.05$ ).

<sup>d</sup> A subpopulation of nuclei in *Agspc72Δ* mutants undergo fast, long-range bypassing (see Figure 2). The numbers in brackets concern these nuclei, which were excluded when the average of parameters was determined.

<sup>e</sup> Each bypassed nucleus was counted.

strains slowly growing colonies with bi- and anucleated cells (Soues and Adams, 1998; Hoepfner *et al.*, 2002; Usui *et al.*, 2003). Based on analysis in budding yeast, it is likely that deletion of *AgSPC72* also prevents anchoring of cMTs at the OP. Previously, we showed that nuclear migration within the multinucleate *A. gossypii* hyphae can be broken down into four types of movement: rotation; forward/backward oscillation and bypassing of nuclei, which are cMT-dependent processes; and cotransport with the cytoplasmic stream, the only observable nuclear movement in the absence of cMTs (Lang *et al.*, 2010). Because of this later microtubule-independent mechanism of tip-directed nuclear movement, it seemed likely that deletion of *AgSPC72* will not be lethal but would affect cMT-dependent nuclear mobility.

Indeed, *Agspc72Δ* mutants were viable and, as shown above, displayed radial colony growth like wild-type even though nuclei were less evenly distributed and formed small clusters (Figure 1, D–F). We monitored the movements of the first five nuclei in four growing hyphae of *Agspc72Δ* for a total of 800 half-min time intervals. Alternating forward/backward movements were not found, and only slow forward movements were observed that rarely exceeded the growth speed of the hyphae (Figure 2, A and B; Table 2; and Supplemental Movies S1 and S2). The one exception was in a rapidly forward moving nucleus able to bypass four nuclei as documented in the image series and the graph labeled *AgH4-GFPspc72Δ(fast)* in Figure 2, A and B (see also Supplemental Movie S3). Further investigation of 70 nuclei in 10 hyphae revealed that three of these highly motile nuclei, which could move with maximum speed of  $8.9 \mu\text{m}$  in 30 s, 3 times faster than the maximum speed in wild type (Table 2). Thus, *Agspc72Δ* hyphae seem to contain two populations of nuclei: a subpopulation of highly mobile nuclei able to undergo long-range bypassing and the majority of nuclei, in which nuclear migration is controlled by cytoplasmic streaming and not by cMTs.

Next, we examined microtubules by immunostaining with antibodies to  $\alpha$ -tubulin and by live cell imaging of hyphae expressing substoichiometric levels of GFP-labeled  $\alpha$ -tubulin (GFP-AgTub1) in addition to wild-type  $\alpha$ -tubulin. Both methods can be used for visualizing cMTs in *A. gossypii* but with some limitations (Gladfelter *et al.*, 2006; Lang *et al.*,

2010). The very thin cMTs of *A. gossypii* are refractory to fixation and are destabilized by more than a substoichiometric concentration of GFP-AgTub1. Compared with wild type, nuclei in *Agspc72Δ* hyphae lack short cMTs; but, surprisingly, a few long cMTs were observed attached or detached from nuclei (Figures 3, A and B, and 7; and Supplemental Movies S7–S9).

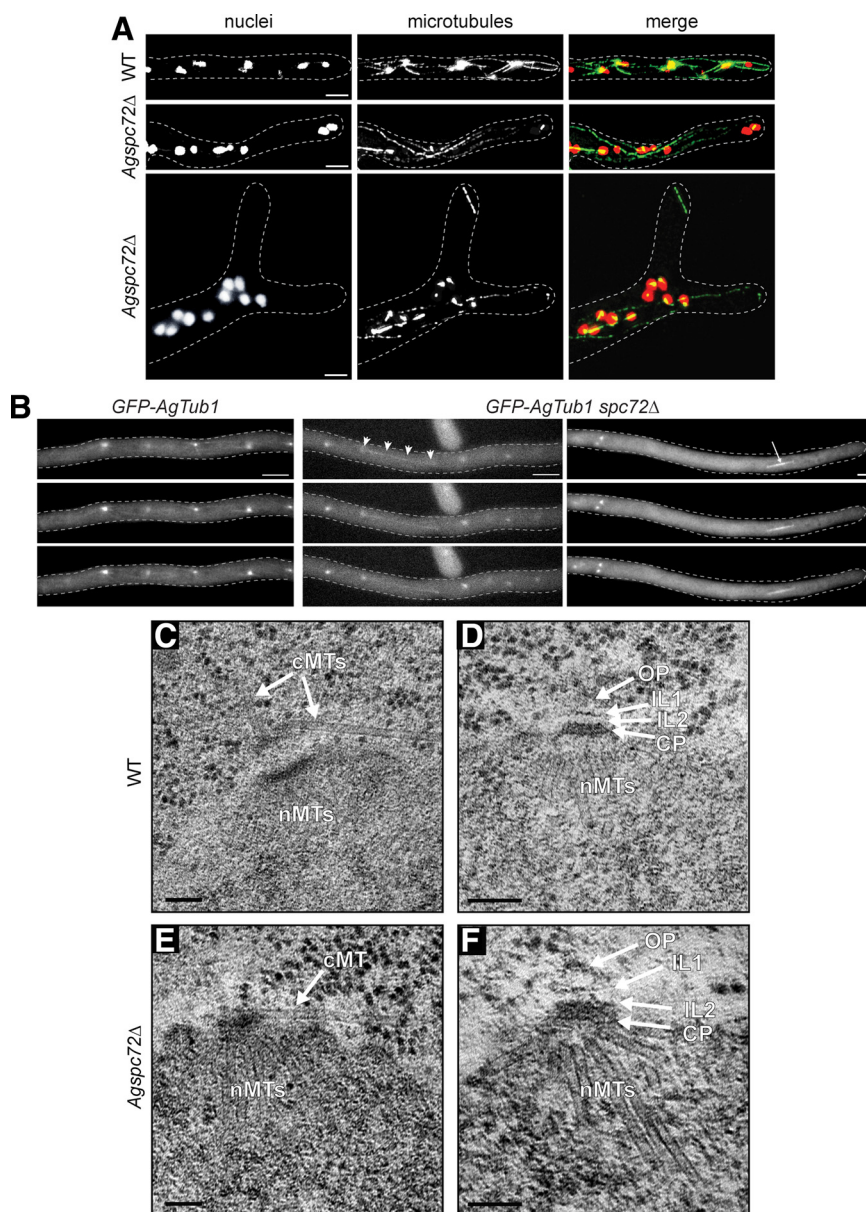
We used serial section EM to analyze the direction in which the few remaining cMTs emanate from *Agspc72Δ* mutant SPBs because cMTs emerge from wild-type SPBs in a perpendicular and a tangential direction (Figure 3C) as described previously (Lang *et al.*, 2010). EM analysis of 33 *Agspc72Δ* mutant SPBs showed no perpendicular cMTs but one or two tangential cMTs attached to the OP region of five mutant SPBs (Figures 3E and 8A). This lack of perpendicular (short) cMTs and the low frequency of tangential (long) cMTs are consistent with our immunofluorescence and live cell imaging data; in *GFP-AgTUB1 spc72Δ* cells, only 31% (12/39) of the SPBs were associated with cMTs, and virtually all of these were long cMTs  $>5 \mu\text{m}$  (Figures 3B and 7 and Supplemental Movies S8 and S9). The few nuclei that contain SPBs attached to long tangential cMTs are probably those that rapidly migrate through the *Agspc72Δ* hyphae, thereby bypassing other nuclei. Together, this indicates that short perpendicular cMTs are probably important for oscillatory movements and long tangential cMTs for nuclear bypassing.

The serial EM sections also revealed structural differences between wild-type and mutant SPBs (Figures 3, D and F, and 8B). A clearly distinguishable OP and IL1 was seen in only 23 and 40% of *Agspc72Δ* SPBs, respectively, compared with 67 and 70% of wild type (*Agspc72Δ*,  $n = 30$ ; wild type,  $n = 46$ ; we excluded SPBs in this analysis if serial section images were not available). Other layers of the SPB were unaffected by deletion of *AgSPC72*.

#### Deletion of *AgNUD1* Results in Loss of the OP but Not Nuclear Dynamics

The budding yeast gene *ScNUD1* encodes an important component of the OP. Deletion of the gene is lethal, probably due to the role of Nud1 as a signaling scaffold for the essential MEN (reviewed in Hoyt, 2000; Pereira and Schiebel, 2001; Stegmeier and Amon, 2004). However, some

**Figure 3.** Nuclear movement, cMTs and SPB structure in *Agspc72Δ*. (A) Wild-type and *Agspc72Δ* mutants were stained with Hoechst to visualize DNA and anti- $\alpha$ -tubulin antibodies to detect microtubules. In the bottom image of *Agspc72Δ*, a cluster of nuclei can be seen at a branch site. These hyphae lack short cMTs, whereas long cMTs that extend along the growth axis are still present. A detached microtubule can be seen in the upper tip region. Bars, 5  $\mu$ m. (B) Representative, deconvolved images of a Z-stack of a wild-type and two *Agspc72Δ* hyphae expressing *GFP-AgTUB1*. The complete stacks are available as Supplemental Movies S7, S8, and S9. Long and short cMTs can be seen emerging from bright foci that represent the SPBs in wild-type, but *Agspc72Δ* SPBs often lack associated cMTs. Arrowheads point to a long cMT in *Agspc72Δ*, which may facilitate bypassing. An arrow indicates an anaphase spindle. Bar, 5  $\mu$ m. (C) EM image of SPB-attached microtubules in wild-type hyphae. Bar, 100 nm. The SPB is associated with nuclear microtubules (nMTs) at the IP and a tangential and perpendicular microtubule at its OP (arrows). (D) SPB structure in wild-type. Bar, 100 nm. A central plaque (CP) plus IL1 and IL2 are marked by arrows. A small amount of amorphous material could also be detected above IL1 and is part of the outer plaque (OP) (Lang *et al.*, 2010). (E) EM image of SPB-attached microtubules in *Agspc72Δ*. Bar, 100 nm. The SPB is associated with nMTs at the IP and a tangential microtubule close to the central plaque (arrow). In this and other thin sections, we never observed perpendicular microtubules. (F) SPB structure in *Agspc72Δ*. Bars, 100 nm. A CP plus IL1 and IL2 are marked by arrows. In some cases, a small amount of amorphous material could also be detected above IL1, which could be part of the OP. However, the size of this OP remnant was substantially reduced compared with a wild-type OP (Lang *et al.*, 2010).



viable point mutants of this gene were characterized and display disturbed organization of cMTs (Gruneberg *et al.*, 2000), but the role of ScNud1 in OP structure could not be determined.

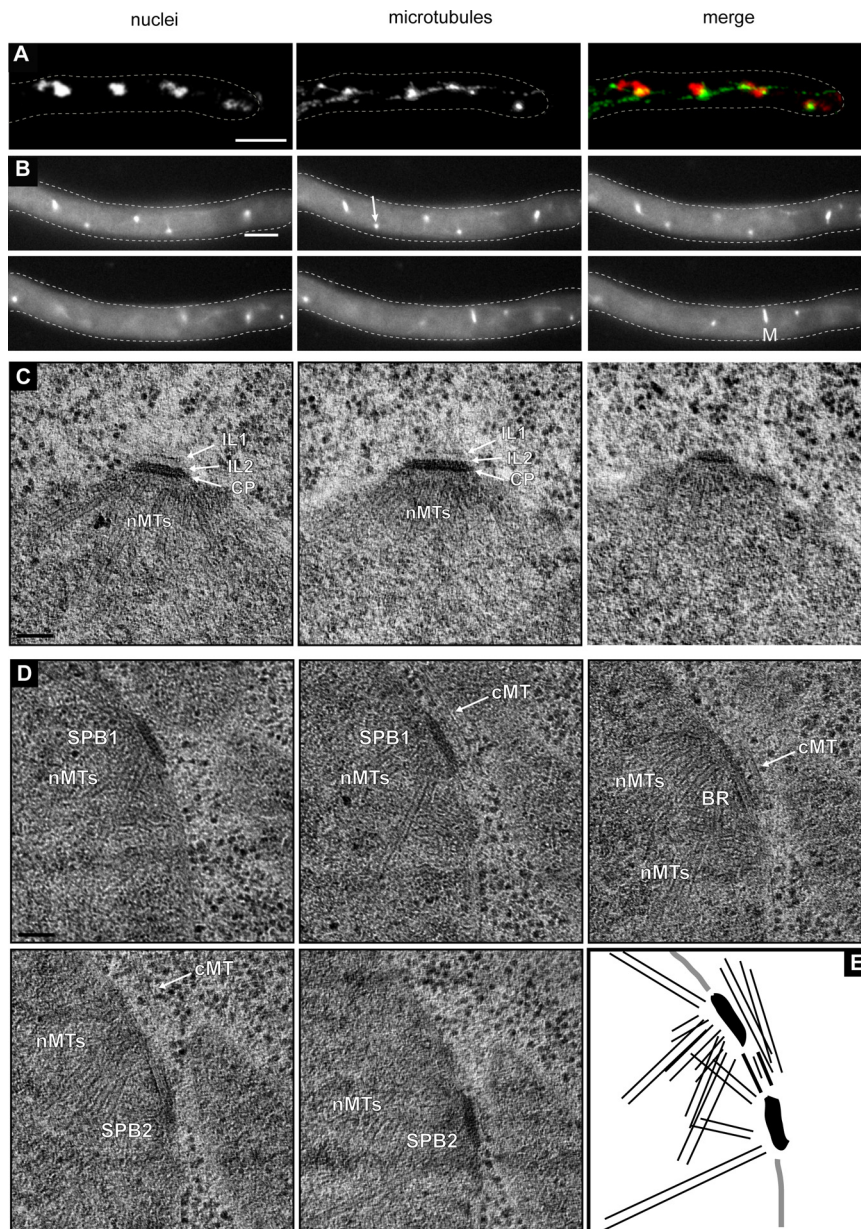
Surprisingly, the *Agnud1Δ* deletion mutant is viable displaying wild-type radial growth on agar (Figure 1D). Nuclear spacing was similar as in wild-type hyphae, although a tendency to form clusters was noted, as concluded from the increased (11%) number of nuclei with <3- $\mu$ m distance to their nearest neighbor (Figure 1F). Forward/backward movements and bypassing of *Agnud1Δ* nuclei were observed as in wild-type (Figure 2, A and B, and Supplemental Movie S4). Examination of 800 half-min time intervals revealed a slight but statistically significant ( $p < 0.01$ ) decrease in nuclear oscillation and a clear decrease in bypassing frequencies compared with wild type (Table 2). Interestingly, *Agnud1Δ* nuclei seemed to have a larger diameter than wild-type or other mutant nuclei we examined (Figure 2A). The reason for the difference is currently un-

known but could be due to a delay in the nuclear division cycle or an increase in ploidy.

Hyphae lacking *AgNUD1* contained both long and short cMTs, and analysis of GFP-AgTub1 images revealed that 89% (41/46) of the *Agnud1Δ* SPBs nucleated both types of cMTs (Figures 4, A and B, and 7; and Supplemental Movie S10). However, very long cMTs >12  $\mu$ m were seldom observed in *Agnud1Δ* mutants. Unlike *Agspc72Δ* mutants, in which cMTs are often detached from nuclei (Figure 3A), we did not detect detached microtubules in the cytoplasm of >20 *Agnud1Δ* hyphae examined.

Analysis of 18 mutant SPBs by EM revealed a complete lack of the OP and a surprising arrangement of SPB-associated cMTs (Figures 4, C-E, and 8, A and B). Serial sections revealed that all duplicated mutant SPBs ( $n = 6$  of 18) had cMTs attached to the bridge structure, approximately half in a perpendicular and half in a tangential direction (Figures 4D and 8A). We therefore conclude that *A. gossypii* SPBs that lack an OP can nucleate and apparently stably anchor cMTs





**Figure 4.** SPBs of *Agnud1Δ* lack an OP and nucleate cMTs from the bridge. (A) *Agnud1Δ* mutants were stained with Hoechst to visualize DNA and anti- $\alpha$ -tubulin antibodies to detect microtubules. Long cMTs that extend along the growth axis are still present and also a few short cMTs were detected. Bar, 5  $\mu$ m. (B) Representative, deconvolved images of a Z-stack from *Agnud1Δ* hypha expressing GFP-*AgTUB1*. The complete stack is available as Supplemental Movie S10. Long and short cMTs can be seen emerging from both sides of a mitotic spindle (M) and from bright foci that represent the SPBs. An arrow points to a SPB lacking cMTs. Bar, 5  $\mu$ m. (C and D) EM images of *Agnud1Δ* mutants. Bars, 100 nm. (C) Serial section images of a single SPB in an *Agnud1Δ* mutant. The central plaque (CP) and IL1 and IL2 were observed, whereas the outer plaque could not be detected in any section. (D) Serial section images of duplicated side-by-side SPBs (SPB1 and SPB2) connected by a bridge (BR) in the *Agnud1Δ* mutant. Nuclear microtubules (nMTs) as well as two cMTs can be seen. The cMTs seem to emerge from the bridge region. (E) Schematic summarizing the five serial section images of D.

from the bridge, presumably through AgSpc72. Furthermore, because nuclear oscillation and bypassing are still observed in *Agnud1Δ* mutants despite a reduced number of cMTs, microtubule anchorage, rather than the number of nucleated cMTs, is probably a key factor for nuclear migration dynamics in *A. gossypii*.

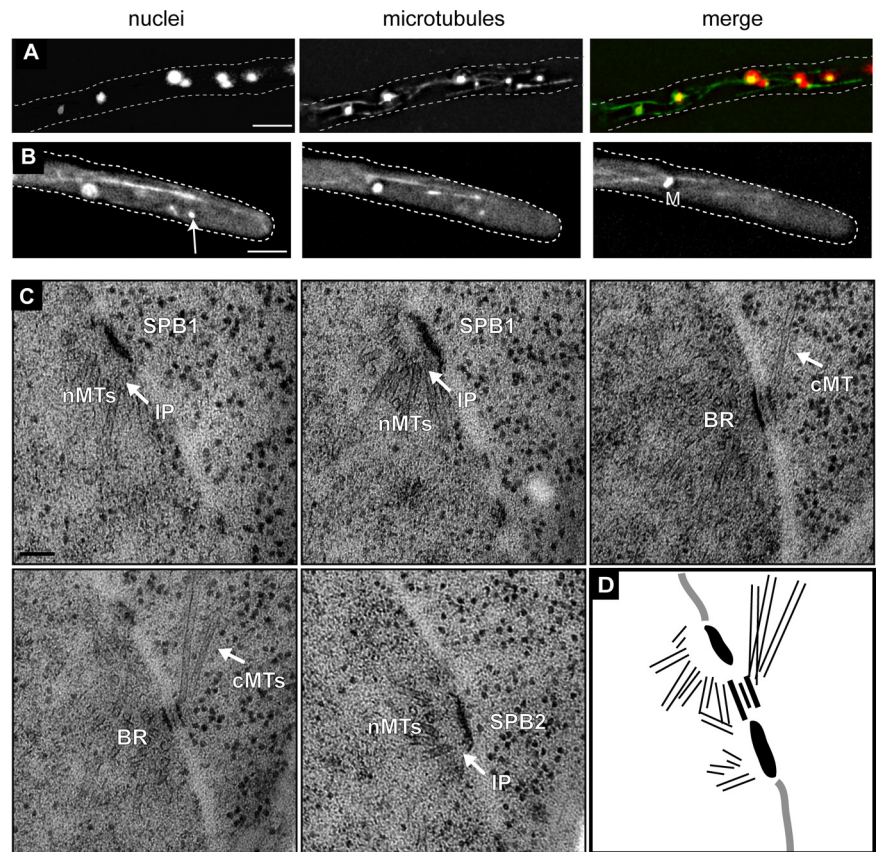
#### Deletion of *AgCnm67* Does Not Impair Nuclear Oscillation or Bypassing

In budding yeast, Cnm67 attaches the OP to IL2, and cells lacking Cnm67 have no OP. This is lethal in some strains or leads to reduced colony growth in other strains because cells with no, two or many nuclei form due to a defect in nuclear positioning at the bud neck before anaphase (Brachat *et al.*, 1998; Hoepfner *et al.*, 2000; Schaerer *et al.*, 2001). Based on these data from budding yeast, we anticipated an *AgCnm67* deletion to be viable but have nuclear migration defects. It is possible that *Agcnm67Δ* mutants would be able to nucleate

cMTs from the bridge/half-bridge as we observed in *Agnud1Δ* and has been observed in cells lacking *ScCnm67* (Brachat *et al.*, 1998).

We found that *Agcnm67Δ* colonies grow like wild-type and that nuclear spacing was only mildly affected (Figure 1, D–G). *Agcnm67Δ* nuclei were still able to undergo oscillation and bypassing (Figure 2, A and B, and Supplemental Movie S5). After nuclear oscillation and bypassing in 20 mutant nuclei for a total of 770 half-min intervals revealed a doubled frequency in bypassing events compared with wild-type, concomitant with increased forward and decreased backward movements (Table 2). Whereas the changed ratio in forward and backward movements, as defined in Table 2, can in part be explained by a slightly faster growth speed of the investigated *Agcnm67Δ* hyphae (Lang *et al.*, 2010), the increased frequency in bypassing is probably a result of changes in cMT organization in this mutant.

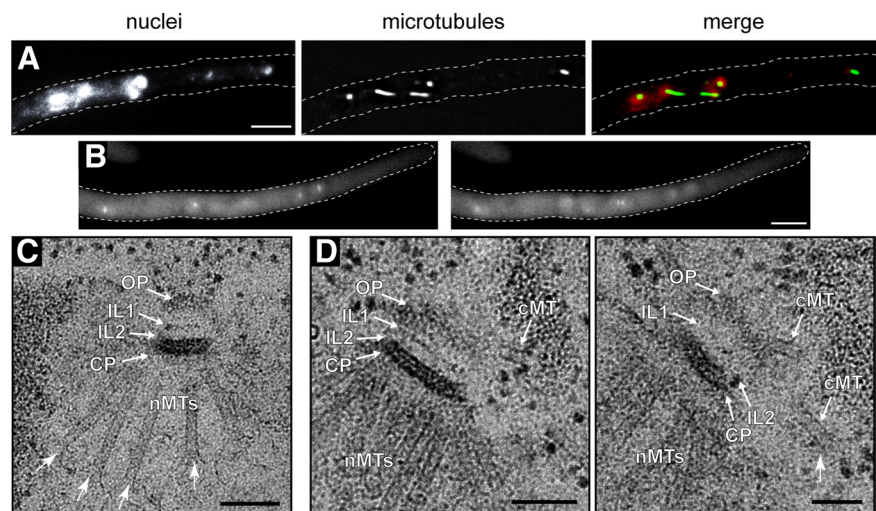




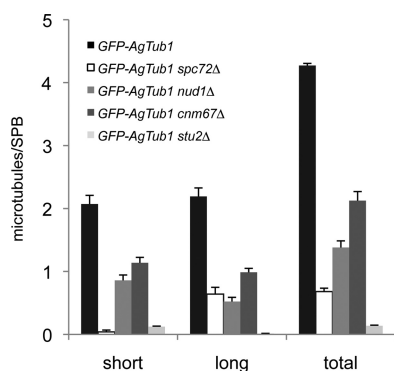
**Figure 5.** cMT nucleation and SPB structure in *Agcnm67* $\Delta$  hyphae. (A) *Agcnm67* $\Delta$  hypha stained with Hoechst to visualize DNA and anti- $\alpha$ -tubulin antibodies to detect microtubules. Long cMTs that extend along the growth axis are present and a few short cMTs were detected. Bar, 5  $\mu$ m. (B) Representative, deconvolved images of a Z-stack of an *Agcnm67* $\Delta$  hypha expressing GFP-AgTub1. The complete stack is available as Supplemental Movie S11. Long and short cMTs can be seen emerging from both sides of a mitotic spindle (M) and from bright foci that represent the SPBs. An arrow points to a SPB lacking cMTs. Bar, 5  $\mu$ m. (C) Serial sections of duplicated side-by-side SPBs (SPB1 and SPB2) connected by a bridge (BR), which nucleates three cMTs in an *Agcnm67* $\Delta$  mutant. Nuclear microtubules (nMTs) are formed at the SPB IP. Images were aligned using AutoAligner to produce the schematic in D. Bars, 100 nm.

Immunostaining with antibodies to  $\alpha$ -tubulin and Z-stacks of GFP-AgTub1 labeled microtubules showed a reduced number of short and long cMTs attached to *Agcnm67* $\Delta$  nuclei compared with wild-type nuclei (Figures 5, A and B, and 7; and Supplemental Movie S11). The Z-stacks also revealed that 78% (72/92) of the *Agcnm67* $\Delta$  SPBs nucleated short, long, and very long (>12  $\mu$ m) cMTs. Examination of 33 mutant SPB structures by EM showed a complete lack of the OP and IL1; but for 11 SPBs, cMTs emanated either from a half-bridge or a bridge structure in both perpendicular and tangential directions (Figures 5, C and D,

and 8, A and B). This confirms that *A. gossypii* SPBs lacking an OP and an IL1 can nucleate and anchor cMTs from a bridge and presumably also half-bridge. It also shows that, in the presence of very long cMTs and a reduced number of short cMTs per SPB, the frequency in nuclear bypassing can increase. It would be interesting to test whether the half-bridge component AgKar1 participates in this rescue mechanism for cMT nucleation in the absence of an OP. But because AgKAR1 is essential (see above), we could not examine the phenotype of *Agcnm67* $\Delta$  *kar1* $\Delta$  or *Agnud1* $\Delta$  *kar1* $\Delta$  double mutants.



**Figure 6.** Microtubule stability and SPB structure in *Agstu2* $\Delta$  mutants. (A) *Agstu2* $\Delta$  hypha stained with Hoechst to visualize DNA and anti- $\alpha$ -tubulin antibodies to detect microtubules. Only short cMTs were detected. Bar, 5  $\mu$ m. (B) Representative, deconvolved images of a Z-stack of an *Agstu2* $\Delta$  hypha expressing GFP-AgTUB1. The complete stack is available as Supplemental Movie S12. (C) EM image of a SPB from *Agstu2* $\Delta$  showing OP, IL1, IL2, and central plaque (CP) layers as well as nuclear microtubules (nMTs). (D) Serial sections of an *Agstu2* $\Delta$  SPB also show the layered SPB structure and nMTs as in A. In addition, three very short cMTs are visible, which have both perpendicular and tangential orientations with the SPB. Arrows point to “flared” microtubule ends frequently seen by EM in *Agstu2* $\Delta$  mutants. Bars, 100 nm.



**Figure 7.** Quantitation of short and long cMTs in hyphae expressing GFP-AgTUB1. The number of short ( $\leq 5\text{-}\mu\text{m}$ ) and long ( $> 5\text{-}\mu\text{m}$ ) cMTs per SPB was quantitated in wild-type and mutant hyphae expressing GFP-AgTUB1, and average values observed for both classes of microtubules are presented along with a combined total. Error bars indicate SE of the mean ( $n = 77, 39, 46, 92,$  and  $58$  SPBs for GFP-AgTUB1, GFP-AgTUB1 *spc72Δ*, GFP-AgTUB1 *nud1Δ*, GFP-AgTUB1 *cnm67Δ*, and GFP-AgTUB1 *stu2Δ*, respectively). Note that only a minor fraction of the long cMTs in GFP-AgTUB1 *spc72Δ* and GFP-AgTUB1 *nud1Δ* are  $> 12\text{ }\mu\text{m}$ , whereas in GFP-AgTUB1 *cnm67Δ* mutants, a substantial fraction is  $> 12\text{ }\mu\text{m}$ .

### AgSTU2 Is Required for Nuclear Oscillation and Bypassing

Although Stu2 is not considered to be a core SPB component in budding yeast in the sense that it is required to maintain the structural integrity of the complex (Jaspersen and Winey, 2004), its localization to the SPB and microtubule plus-ends, biochemical and genetic interactions with SPB OP and  $\gamma$ -tubulin components, microtubule binding domain, and regulation of microtubule dynamics made it a leading candidate to participate in cMT organization (Wang and Huffaker, 1997; Chen *et al.*, 1998; Kosco *et al.*, 2001; Severin *et al.*, 2001; Pearson *et al.*, 2003; Usui *et al.*, 2003; van Breugel *et al.*, 2003; Al-Bassam *et al.*, 2006), and thus control nuclear migration dynamics in *A. gossypii*. The *A. gossypii* genome contains a single XMAP215-like gene, AgSTU2, that is a syntenic homologue of *S. cerevisiae* STU2. AgStu2 and ScStu2 are 32% identical.

If AgStu2 is needed for cMT growth or stability, then we would expect to see short or absent microtubules in *Agstu2Δ*, which should manifest itself in defects in cMT-dependent forms of nuclear migration. Surprisingly, deletion of AgSTU2 resulted in viable cells that grew on agar plates similar to wild-type (Figure 1C). Labeling of *Agstu2Δ* nuclei with H4-GFP showed that nuclear spacing is affected: hyphae displayed enlarged distances from the leading nucleus to the tip region as well as decreased distances between adjacent nuclei (Figure 1, E–G). Nuclei in *Agstu2Δ* hyphae only migrated forward with the cytoplasmic stream (Figure 2, A and B, and Supplemental Movie S6). In 200 half-min intervals, no backward movement and no nuclear bypassing were observed (Table 2), indicating a lack of cMTs.

Analysis of microtubules by immunostaining with anti- $\alpha$ -tubulin antibodies showed some *Agstu2Δ* nuclei associated with very short cMTs, whereas long cMTs were absent (Figure 6A). Quantitation of microtubule length from immunofluorescence images revealed an average cMT length of  $200 \pm 56\text{ nm}$  ( $n = 31$  cMTs/51 SPBs) in *Agstu2Δ* mutants compared with  $1912 \pm 93\text{ nm}$  ( $n = 275$  cMTs/75 SPBs) in wild-type hyphae (data not shown). We also examined microtubules by live cell imaging using GFP-AgTub1 and found that cMTs were virtually undetectable (Figures 6B

and 7 and Supplemental Movie S12). It is likely that GFP-AgTub1 leads to slightly less stable microtubules *in vivo*, which when combined with *Agstu2Δ*, further decreases cMT stability. Thus, the cMT number and length observed by immunofluorescence is very likely correct.

Unlike the other mutant SPBs we examined by EM, all known SPB layers were observed in *Agstu2Δ* and their dimensions are very similar to wild-type SPBs (Figures 6, C and D, and 8B). Even though the OP structure seems to be preserved, serial section EM analysis confirmed the significant reduction in the number of cMTs nucleated at *Agstu2Δ* SPBs: only 39% of the mutant OPs ( $n = 7$  of 18) compared with 98% wild-type ( $n = 49$  of 50) SPBs were associated with cMTs (Figure 8A). The cMTs we observed by EM were rather short but showed both perpendicular and tangential association to the SPB (Figures 6D and 8A). We also noticed that most of the non-SPB-associated microtubule ends in *Agstu2Δ* seemed to exhibit a “flared” or “peeling” structure, which has been observed previously in other EM studies and is indicative of shortening microtubules (Byers *et al.*, 1978; Mandelkow *et al.*, 1991; O’Toole *et al.*, 1999). This phenotype was rarely detected in wild-type cells or other mutants we examined. From our analysis of *Agstu2Δ*, we conclude that cMTs nucleated in this mutant are too short to induce nuclear oscillations or even bypassing. The fact that *Agstu2Δ* is able to form both perpendicular and tangential attachments even though we never observed cMTs  $> 200\text{ nm}$  in this mutant (Figure 8, A and B;  $n = 60$  cMTs) suggests that something other than microtubule length determines the orientation of cMTs. The short cMTs and normal SPB structure suggest that AgStu2 functions in plus-end microtubule dynamics and does not have a structural role at the SPB.

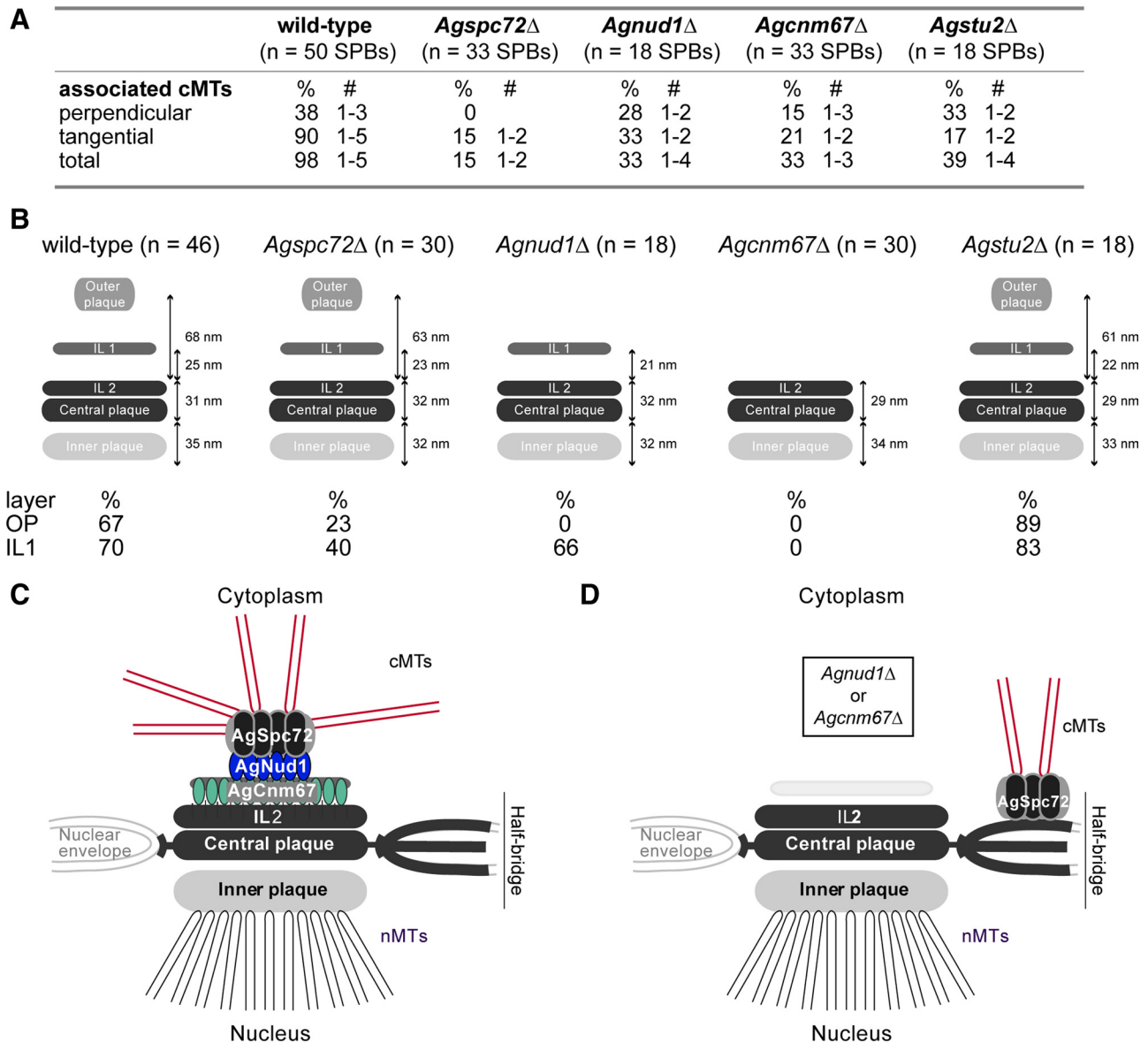
## DISCUSSION

From our EM analysis of SPB structure in *Agspc72Δ*, *Agnud1Δ* and *Agcnm67Δ* mutants, we propose that the *A. gossypii* organelle is assembled in a stepwise manner. By analogy to *S. cerevisiae* SPB assembly, binding of AgCnm67 drives assembly of AgNud1, followed by AgSpc72 and the  $\gamma$ -tubulin complex to form the OP (Figure 8C). In wild-type hyphae, an intact OP is essential for nucleation and anchorage of cMTs involved in nuclear oscillations and bypassing. AgStu2 also plays an important role in the formation of cMTs and thus nuclear movements, although it is probably not involved in the structural maintenance of the SPB.

### Role of cMTs in Nuclear Oscillations and Bypassing

A major goal of this work was to test the model for cMT-dependent nuclear movements proposed in our previous publication (Lang *et al.*, 2010). Shown schematically in Figure 9, we proposed that arrays of short perpendicular and long tangential cMTs emanate from nuclear SPBs in the multinucleate hyphae of *A. gossypii*. Each of the three nuclei shown in the wild-type hypha is associated with short and long cMTs growing in apical and subapical directions from its OP. The most apical nucleus makes close contact to the growing tip via several cMTs. The other two nuclei are connected with the hyphal cortex via short cMTs and, based on preliminary data obtained with plus-end-labeled cMTs, also via their long cMTs (Lang *et al.*, 2010; Grava and Philippsen, unpublished observations). We hypothesized that growth and shrinkage of cMTs provides pulling and pushing forces for short-range nuclear oscillations and that long-range movements during nuclear bypassing are probably achieved by pulling forces of the long cMTs when the cortex connection of short cMTs is reduced or absent. The four dele-





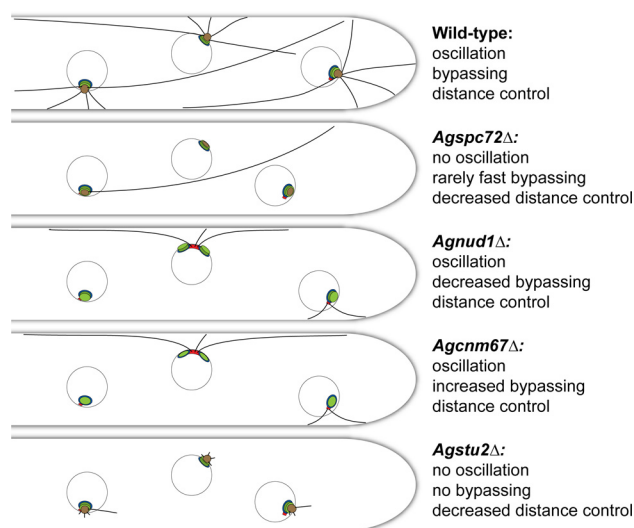
**Figure 8.** Comparison of *A. gossypii* wild-type and mutant SPB structure and cMT nucleation based on EM analysis. (A) Quantitation of cMTs that emerge per SPB in wild-type and mutants based on EM analysis. (B) Schematics of wild-type and mutant SPBs depicting the observed layers and averaged distances between the layers, based on measurements shown in Supplemental Table S3. The percentage of SPBs in which the OP and IL1 were seen is indicated below. (C) Model of the *A. gossypii* SPB based on SPB morphology we observed in different deletion mutants. The layered SPB of *A. gossypii* is composed of AgCnm67 in IL1, which recruits AgNud1. Next, AgSpc72 binds to form the OP and tether the  $\gamma$ -tubulin complex that nucleates cMTs. (D) In the absence of AgCnm67 or AgNud1, cMTs are formed from the bridge region (Brachat *et al.*, 1998; Gruneberg *et al.*, 2000; Hoepfner *et al.*, 2000). Studies in budding yeast lead us to hypothesize that AgSpc72 and the  $\gamma$ -tubulin complex relocate here (Knop and Schiebel, 1998), at least in these mutants. In *Agnud1* $\Delta$ , IL1 remains, whereas it is lost in *Agcnm67* $\Delta$ .

tion mutants that we have analyzed here each have unique alterations in their cMT arrays compared with wild-type (summarized in Figures 7 and 8) that allowed us to test various aspects of this model. Based on the changes in cMT organization that we observed, we can explain most of the corresponding changes in nuclear mobility and in distance control in the mutants. In addition, although the *A. gossypii* SPB structure and core proteins are similar to *S. cerevisiae*, our analysis of the deletion mutants revealed unanticipated phenotypes that differed considerably from those described previously in budding yeast (Rose and Fink, 1987; Geissler *et al.*, 1996; Spang *et al.*, 1996; Knop

*et al.*, 1997; Brachat *et al.*, 1998; Chen *et al.*, 1998; Pereira *et al.*, 1999; Gruneberg *et al.*, 2000; Hoepfner *et al.*, 2000, 2002; Usui *et al.*, 2003), shedding light on the mechanism of cMT anchorage, SPB assembly, and the evolution of conserved cytoskeletal proteins in multinucleated hyphae.

#### Formation of the OP and cMT Nucleation: AgSpc72

In *Agspc72* $\Delta$  mutants, the majority of nuclei do not nucleate cMTs, therefore, they lack cMT-dependent movements and migrate with the cytoplasmic stream (Figure 9). The lack of cMTs in *Agspc72* $\Delta$  is similar to *Scspc72* $\Delta$  mutants, which form only very short and unstable cMTs (Soues and Adams,



**Figure 9.** Model of the cytoplasmic sides of SPBs and attached cMTs in wild-type and mutant hyphae of *A. gossypii*. The four deletion mutants presented here have altered cMT arrays compared with wild-type, and these alterations can explain most of the observed changes in nuclear mobility and in distance control to the hyphal tip assuming that microtubules can exert pulling forces upon cortical contacts, which may be transient, e.g., causing oscillations, or more stable, e.g., for nuclear bypassing driven by long microtubules. See Discussion for details.

1998; Hoepfner *et al.*, 2002). In a few rare instances, we observed long tangential microtubules and an associated OP by EM in *Agspc72Δ* mutants; it seems that the long cMT can pull the attached nucleus over long distances, thereby rapidly bypassing several nuclei. The fact that short cortex-associated cMTs that may interfere with bypassing are also absent in *Agspc72Δ* hyphae could explain the high speed of these rare bypassing events. The lack of short cMTs also accounts for the increased distance of the most apical nucleus to the tip.

However, the question of how any cMTs are nucleated and attached at such a mutant SPB is unclear because *Agspc72Δ* mutants are lacking the presumed cytoplasmic anchor for the  $\gamma$ -tubulin complex. Due to the fact that budding yeast Stu2 binds to Spc72 and the  $\gamma$ -tubulin complex through sequences in its N terminus and stimulates polymerization of pure tubulin *in vitro* (Usui *et al.*, 2003; van Breugel *et al.*, 2003; Al-Bassam *et al.*, 2006), AgStu2 was a candidate to help anchor cMTs to the SPB in the absence of AgSpc72, but our experimental data do not support this hypothesis. Another candidate for anchoring the  $\gamma$ -tubulin complex at the OP in the absence of AgSpc72 is AgSpc110. In *S. cerevisiae*, Spc110 tethers  $\gamma$ -tubulin to the SPB IP but it can function in cMT nucleation if it is artificially tethered to the OP (Knop and Schiebel, 1998). Normally, Spc110 is targeted to the nucleus by an N-terminal nuclear localization sequence (Adams and Kilmartin, 1999). This domain is not conserved in AgSpc110, and other nuclear localization motifs were not detected (Lang and Jaspersen, unpublished observations), so how AgSpc110 enters the nucleus is unknown. A portion of AgSpc110 might remain in the cytoplasm. The *A. gossypii* genome also carries multiple *no* homologs in *bakers yeast* (NOHBY) genes that could play a role in SPB functions specific to multinucleate fungi, including nucleation of different classes of cMTs (Brachat *et al.*, 2003).

### Loss of AgNUD1 and AgCNM67 Reveals a Conserved Rescue Mechanism for cMT Nucleation at the SPB

In comparison with hyphae lacking AgSpc72, the absence of AgCnm67 or AgNud1 results in a radical change in SPB structure and microtubule organization: the OP is lost and cMTs are nucleated from the bridge/half bridge (Figure 8D). However, nuclear oscillation and bypassing are still observed in both mutants, indicating that bridge/half-bridge-nucleated cMTs are sufficient to direct typical nuclear movements within hyphae (Figure 9). In both mutants the total number of cMTs per nucleus is reduced approximately two-fold compared with wild-type; therefore, anchorage to the SPB rather than the total number of nucleated cMTs seems to be a critical factor controlling both nuclear oscillations and bypassing. The increased frequency in bypassing we observed in *Agcnm67Δ* mutants compared with wild-type might be due to the relatively high abundance of very long cMTs concomitant with an overall decreased number of cMTs, which could physically obstruct rapid long-range movement of a large nucleus or reduce the tethering force that resists bypassing. Due to the larger size of *Agnud1Δ* nuclei, their movement is probably more restricted so a decrease in bypassing was observed.

Nucleation of microtubules from the half-bridge/bridge seems to be an evolutionary conserved rescue mechanism in the absence of an OP because it is also observed in budding yeast *cnm67Δ* and *nud1* mutants (Brachat *et al.*, 1998; Adams and Kilmartin, 1999; Gruneberg *et al.*, 2000). Nucleation of microtubules from the half-bridge also occurs in wild-type budding yeast cells during G1 phase of the cell cycle (Byers and Goetsch, 1975). Although half-bridge microtubules are rarely observed in asynchronous cultures of *S. cerevisiae*, they can be easily visualized in cells treated with the mating pheromone  $\alpha$ -factor because this class of cMTs is essential for the nuclear migration and fusion after mating (karyogamy). During our EM analysis of *A. gossypii* wild-type SPBs, we could not find evidence of cMT nucleation from the half-bridge simply because this structure was only observed twice (Lang *et al.*, 2010). *A. gossypii* is not known to undergo nuclear fusions, so half-bridge-emanating microtubules may have been lost during evolution.

Perhaps one of our most unexpected findings was that *Agnud1Δ* mutants are viable, because most *nud1* mutants in *S. cerevisiae* arrest cell division in anaphase due to the role of Nud1 in MEN at the SPB OP (reviewed in Hoyt, 2000; Pereira and Schiebel, 2001; Stegmeier and Amon, 2004). Our finding that AgNUD1 is not essential suggests that the MEN is either nonessential in *A. gossypii* or does not depend on SPB recruitment for its activation. Due to the growth mode of *A. gossypii*, in which segregation of chromosomes, an SPB, and other cellular material into the daughter cell is not critical for survival, we suspect that positioning of the nucleus and spindle is probably not under tight checkpoint control as it is in *S. cerevisiae*. Indeed, preliminary experiments with some *A. gossypii* MEN genes show that they are not essential (Finlayson and Philippsen, unpublished observations). Although Nud1 is one of the few core SPB components with recognizable homologues in other species, the functional conservation among these proteins is poor, aside from their role in astral microtubule formation. Given our findings, Nud1 orthologues may have a rapidly evolving activity during cell division.

### Regulation of cMTs by AgStu2

Of all the deletions we analyzed, *Agstu2Δ* has the greatest effect on cMT length, which explains the complete lack of



nuclear oscillations and bypassing in this mutant and the 2.5-fold increase in the distance from the tip to the first nucleus compared with wild-type. Although *Agspc72Δ* and *Agstu2Δ* mutants are similar in terms of their effects on nuclear dynamics, the mechanisms responsible for cMT loss in *Agstu2Δ* and *Agspc72Δ* mutants are very different, as shown by our EM analysis and depicted in Figure 9. *AgSpc72* is most likely a structural component of the SPB; therefore, its elimination results in changes in SPB morphology and cMT nucleation, whereas *AgStu2* is a microtubule-associated protein, which affects MT formation/elongation but not SPB structure. Unlike *Agspc72Δ* nuclei that are occasionally able to form very long cMTs and undergo rapid bypassing, we did not observe any bypassing in *Agstu2Δ* hyphae because no long cMTs are able to form. The number of short cMTs was also dramatically reduced, but we were able to detect these forming in both perpendicular and tangential orientations. Most likely, *AgStu2* affects cMT polymerization as it does in many other eukaryotes, including budding and fission yeast, *Aspergillus nidulans*, and metazoans (reviewed in Gard *et al.*, 2003). Curiously, neither nuclear migration nor positioning was affected in *A. nidulans* hyphae lacking the single XMAP215/Stu2 orthologue *alpA*, despite a dramatic reduction in the number and dynamics of cMTs (Enke *et al.*, 2007). Perhaps nuclear dynamics is controlled through different mechanisms in filamentous fungi that use a single MTOC like *A. gossypii* versus multiple MTOCs like *A. nidulans* (Konzack *et al.*, 2005; Lang *et al.*, 2010).

We were also surprised by the fact that deletion of *AgSTU2* did not result in loss of viability as it does in budding yeast. *ScSTU2* is essential due to its role in nuclear microtubule dynamics and bipolar spindle assembly; it also has a nonessential function in cMT polymerization and nuclear positioning (Wang and Huffaker, 1997; Chen *et al.*, 1998; Kosco *et al.*, 2001; Severin *et al.*, 2001; Pearson *et al.*, 2003; Usui *et al.*, 2003; van Breugel *et al.*, 2003; Al-Bassam *et al.*, 2006). It is unclear why *AgSTU2* is not required for mitotic spindle formation as it is in budding yeast. Alterations in cell cycle regulation, the mechanism of spindle assembly, or both could account for this difference.

## CONCLUSIONS

In conclusion, the multinucleate growth mode of *A. gossypii* has resulted in different demands on the microtubule cytoskeleton that have driven evolution of SPB components to fit its unique life-style. Because nuclear migration into a bud once every cell cycle does not apply in *A. gossypii* hyphae, cMTs and SPB OP components are not critical for cell viability but do function in nuclear oscillations and bypassing. One newly evolved cytoskeletal element are tangential long cMTs, which drive nuclear bypassing, whereas the conventional short cMTs, which make contacts with the cell cortex, are important for nuclear oscillations as summarized in Figure 9. Our functional analysis of *A. gossypii* SPB genes has allowed us to better understand the adaptive properties of the cytoskeleton involved in growth and development of many eukaryotic cells.

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