Regulation of Distinct Septin Rings in a Single Cell by Elm1p and Gin4p Kinases

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Septins are conserved, GTP-binding proteins that assemble into higher order structures, including filaments and rings with varied cellular functions. Using four-dimensional quantitative fluorescence microscopy of *Ashbya gossypii* **fungal cells, we show that septins can assemble into morphologically distinct classes of rings that vary in dimensions, intensities, and positions within a single cell. Notably, these different classes coexist and persist for extended times, similar in appearance and behavior to septins in mammalian neurons and cultured cells. We demonstrate that new septin proteins can add through time to assembled rings, indicating that septins may continue to polymerize during ring maturation. Different classes of rings do not arise from the presence or absence of specific septin subunits and ring maintenance does not require the actin and microtubule cytoskeletons. Instead, morphological and behavioral differences in the rings require the Elm1p and Gin4p kinases. This work demonstrates that distinct higher order septin structures form within one cell because of the action of specific kinases.**

INTRODUCTION

Septins are conserved guanosine triphosphate (GTP)-binding proteins that assemble into heteromeric, nonpolar polymers with varied cellular functions (Versele and Thorner, 2005; Weirich *et al*., 2008). Initially discovered in budding yeast based on their role in cytokinesis, septins have been shown to act as barriers to membrane diffusion, as scaffolds for proteins that regulate the cell cycle and cytoskeleton, and as players in cell migration (Hartwell, 1971; Barral *et al*., 2000; Takizawa *et al*., 2000; Gladfelter *et al*., 2001; Finger *et al*., 2003). Septin family members have been found in fungal, animal, and some algal genomes but not in plants (Pan *et al*., 2007). In all cases studied, septin function requires the assembly of septin subunits into higher order complexes in cells. Importantly, septin mutations or aberrant septin expression has been associated with a variety of neurodegenerative diseases as well as some cancers (Ihara *et al*., 2003; Hall and Russell, 2004; Hall *et al*., 2005; Russell and Hall, 2005; Ihara *et al*., 2007; Sitz *et al*., 2008).

Recently, oligomeric septin complexes were observed at the supramolecular level by electron microscopy for budding yeast, worm, and human in vitro-expressed complexes and at the atomic level by x-ray crystallography for three human septins (John *et al*., 2007; Sirajuddin *et al*., 2007; Bertin *et al*., 2008). All studies found that septins form symmetrical or apolar rods of repeating oligomers of septin subunits. The dimensions of the core rod varied depending upon the number of septins coexpressed. These rods can associate end-toend to form nonpolar, striated filaments that are frequently paired in vitro (Frazier *et al*., 1998; Versele *et al*., 2004).

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However, very little is known about how the filaments seen in vitro relate to higher order septin structures observed by either light or electron microscopy in vivo. Septin complexes assemble into diverse higher order structures in cells, including filaments, rings, and "gauzes" of enmeshed filaments (Haarer and Pringle, 1987; Kinoshita *et al*., 2002; Rodal *et al*., 2005). There is still sizeable ambiguity regarding the basis for filament polymerization and how filaments may come together to build such varied higher order structures in vivo.

In budding yeast, five septins (Cdc3p, Cdc10p, Cdc11p, Cdc12p, and Shs1p/Sep7p) colocalize first as a patch and then form an hourglass-shaped collar between the mother and growing bud. At cytokinesis, the ring splits and then disassembles in both the daughter and mother cell (Caviston *et al*., 2003; Dobbelaere *et al*., 2003). Septin ring reorganizations are at least partially controlled by posttranslational modifications of the septin subunits that alter the subunit dynamics and appearance of higher order structures (Johnson and Blobel, 1999; Mortensen *et al*., 2002; Dobbelaere *et al*., 2003; Versele and Thorner, 2004; Egelhofer *et al*., 2008). How specific posttranslational modifications lead to changes in the polymerization of septin filaments or assembly of higher order structures in vivo is not well understood. At the end of each cell cycle, yeast septins are stripped of these modifications and the subunits are reused in subsequent cycles (McMurray and Thorner, 2008). In contrast, in differentiated mammalian cells such as neurons, higher order septin structures persist over extended times. Thus, the yeast model system is likely insufficient to identify mechanisms relating to long-term maintenance and maturation of the septin cortex because the structures are disassembled with the turning of each cell cycle.

We have established a model fungal system, *Ashbya gossypii*, for the study of higher order septin structures in vivo that complements work done both in other fungi and mammalian cells (Dietrich *et al*., 2004; Wendland and Walther,

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With the exception of plasmidic strains (brackets), all analyzed mycelia were homokaryotic (all nuclei have same genotype).

2005; Helfer and Gladfelter, 2006). *A*. *gossypii* is a filamentous ascomycete that diverged at least 100 million years ago from a common ancestor with *Saccharomyces cerevisiae* (Dietrich *et al*., 2004). Notably, there is no uninucleate yeast form in the fungus and instead it grows exclusively as multinucleate hyphae that contain tens of nuclei sharing a common cytoplasm. This multinucleate state arises from asynchronous nuclear division without cell division (Gladfelter, 2006b; Gladfelter *et al*., 2006). In the genome, *A*. *gossypii* has the same set of septin genes as *S*. *cerevisiae* but with an additional, tandem duplication of *CDC11*. In *A*. *gossypii*, the septins are essential to establish septation sites (which form in response to stress and senescence), to promote mitosis, and to form asexual spores (Gladfelter, 2006a; Helfer and Gladfelter, 2006). Importantly, septins are not essential in *A*. *gossypii*. Therefore, we can perturb the structures and still monitor them in living cells. Finally, we are able to measure maturation of the septin cortex in the absence of cell cycle driven remodeling (Helfer and Gladfelter, 2006).

We use four-dimensional (4D) quantitative fluorescence microscopy to show that *A*. *gossypii* septins assemble into morphologically distinct classes of rings that vary in dimensions, intensities, lifetimes, and positions across a single cell. In these studies, we demonstrate that highly variable septin rings coexist in one cell and require different signaling pathways for their assembly and maintenance.

MATERIALS AND METHODS

Growth Conditions and Strain Construction

A. *gossypii* media, culturing, and transformation protocols are described in Wendland *et al*. (2000) and Ayad-Durieux *et al*. (2000) Strains generated and used in this study can be found in Table 1. All DNA manipulations were carried out according to Sambrook and Russell (2001). Plasmids used here are listed in Table 2. Polymerase chain reaction (PCR) was performed using standard methods with polymerases from Roche Diagnostics (Indianapolis, IN) and Invitrogen (Carlsbad, CA). Oligonucleotides (oligos) were synthesized by Integrated DNA Technologies (Coralville, IA), and all restriction enzymes came from New England Biolabs (Beverly, MA). Oligonucleotide

primers are listed in Table 3. Plasmid isolation from yeast and sequencing were performed as described in Schmitz *et al*. (2006). All sequencing was carried out by Dartmouth College Core Facilities (Hanover, NH).

A. *gossypii* deletion mutants were generated using PCR-based, one-step gene targeting approach with dominant selection markers (McElver and Weber, 1992; Baudin *et al*., 1993; Wach, 1996; Wendland *et al*., 2000). For G418 resistance, the deletion cassettes were amplified off the pAGT140 template by using "gene name"-D1/D2 (or F/R for AGO174 and AGO175 only) primer pairs that had \sim 45-bp homology to the termini of the gene of interest and 20-bp homology to the termini of a selection marker. For single deletion mutant strains, correct integration of selection markers into and removal of the gene of interest from the genome was verified by analytical PCR using combinations of oligonucleotides gene name-V1, V2, IV1, IV2, Int (gene) locus-specific sequences), VG3, VG5 (G418-specific sequences), and V2 NAT1 and V3 NAT1 (NAT1-specific sequences). Transformation of multinucleate mycelium leads to heterokaryotic cells containing a mixture of transformed and wild-type nuclei which may not display an apparent phenotype. For subsequent analysis, homokaryotic mycelia were obtained by isolating and growing single spores. To exclude phenotypes because of random mutation, at least two independent transformants were characterized for each strain.

To create strains AG231 (*CDC11a-mCherry-NAT1*) and AG245 (*CDC3-CFP-NAT1*), plasmids containing *CDC11a* and *CDC3* (AGB125 and AGB127) were kindly provided by the Philippsen laboratory (University of Basel, Basel, Switzerland). Oligos AGO299, 190, 192, and 188 containing sequence homology to the 3' of a septin open reading frame (ORF) were used to amplify either *mCherry-NAT1* from AGB048 or *CFP-NAT1* from AGB011. Yeast cotransformation of PCR products with plasmids AGB125 or 127 yielded AGB141 (*CDC11a-mCherry-NAT1*) and AGB100 (*CDC3-CFP-NAT1*). AGB141 was verified by restriction enzyme digest with SalI. AGB100 was verified by digestion with EcoRI and NdeI and sequenced with oligos AGO199, 200, 201, and 202. *A*. *gossypii* cells were transformed with AGB141 to make AG231. Strain AG232 (*CDC11a-mCherry-NAT1*, *SEP7-GFP-GEN3*) was created by digesting plasmid AGB088 (*SEP7-GFP*) with SacI and transforming this product into strain AG231. Verification of these strains was performed using fluorescence light microscopy.

To create strains AG306 (*CDC11a-mCherry-NAT1*, *[pCDC10-YFP-GEN3]*) and AG308 (*CDC11a-mCherry-NAT1*, *[pCDC12-Venus-GEN3]*), *CDC10* and *CDC12* were amplified along with 500 base pairs upstream and downstream of the ORFs out of the *A*. *gossypii* genome by using primer pairs AGO109/110 and AGO115/116. The products were ligated, using T4 ligase (Invitrogen), into pCR 2.1 vectors (Invitrogen) that had been cut with AflII by using TA Cloning kits (Invitrogen) to create plasmids AGB139 (*CDC10*) and AGB122 (*CDC12*). AGB139 was verified by restriction enzyme digest with BglII and HinDIII, and this product was sequenced using oligos AGO107, 108, 111, and 112. AGB122 was verified by digestion with EcoRI and NdeI and sequencing using oligos AG-O-119, 120, 121, and 122. AGB139 and AGB122 were digested with BamH1 and NotI and ligated into pRS416 linearized with BamHI and NotI to create plasmids AGB112 (*CDC10*) and AGB123 (*CDC12*). AGB112 was verified by digestion with NdeI and PvuII. AGB123 was verified by digestion with ScaI and XhoI. A PCR fragment that had homology to *CDC10* and contained *YFP-GEN3* was amplified from plasmid AGB023 (*YFP-GEN3*) by using oligos AGO193 and 198. Yeast cotransformation of this product and AGB112 created AGB098 (*CDC10-YFP-GEN3*). AGB098 was verified by digestion with XmaI and sequenced with oligos AGO202, 206, 207, and 208. This plasmid was then transformed into strain AG127 (*SEP7-GFP-NAT1*) to create strain AG-306. A PCR product with homology to *CDC12* and containing *Venus-GEN3* was amplified from plasmid AGB024 (*Venus-GEN3*) using oligos AGO195 and 197. This product and plasmid AGB123 were cotransformed into yeast to create plasmid AGB136 (*CDC12-Venus-GEN3*). AGB136 was verified by digestion with BamH1 and NdeI and sequencing using oligos AGO202, 206, 208, and 209. AGB136 was transformed into strain AG127 to create strain AG308. Verification of these strains was performed using fluorescence light microscopy.

Colony radial growth assays were performed by spotting equal volumes of spores from each mutant strain beside an isogenic strain carrying the Sep7-GFP-NAT1 construct on selective media. Measurements for four independent experiments for each genotype were collected each day for 10 d of growth at 30°C.

Microscope Setup and Image Processing

An AxioImager-M1 upright light microscope (Carl Zeiss, Jena, Germany) equipped with the following Zeiss oil immersion objectives was used: EC Plan-Neofluar $40\times$ /1.3 numerical aperture (NA), Plan-Apochromat $63\times$ /1.4 NA, Plan-Neofluar $100\times/1.3$ NA, and α Plan-Fluar $100\times/1.45$ NA. For visualization of green fluorescent protein (GFP), filter set 41025 (Chroma Technology, Brattleboro, VT) and filter set 38HE (Carl Zeiss) were used. For visualization of AlexaFluor 568, Chroma filter sets 41002B and 41043 were used. For visualization of Hoechst, Zeiss filter set 49 was used. An Exfo X-Cite 120 lamp was used as the fluorescent light source. Images were acquired with an Orca-AG (C4742-80-12AG; Hamamatsu, Bridgewater, NJ) charge-coupled device camera driven by either OpenLab 5 (Improvision, Lexington, MA) or Volocity 4 (Improvision). Z-stacks of still images were acquired at different slice sizes (0.2–0.5 μ m) and processed by fast or iterative deconvolution using

calculated point spread functions. OpenLab 5 was used to acquire time-lapse movies that were then processed in Volocity 4. All still images were linearly contrast enhanced in Volocity 4 and Photoshop CS2 (Adobe Systems, Mountain View, CA). All images and movies presented are maximum projections of three-dimensional volumes.

Time-lapse Analysis

All time-lapse movies presented here were acquired using the EC Plan-Neofluar $40\times/1.3$ NA objective, 10–20% fluorescence light transmission, exposed for 100 ms per z-step, through 14 1- μ m steps, and then iteratively deconvolved identically using a calculated point spread function (45 iterations). Cells used in time-lapse experiments were germinated and grown on 12–25% Ashbya Full Medium (AFM) 1–3% dextrose 1% agarose gel pads in a humid chamber for \sim 13 h at 30°C and imaged in a humid chamber at room temperature $(\sim 23^{\circ}\text{C})$. Volocity 4 was used to measure ring lengths and intensities. Ring length was defined as the distance between the visible (by Sep7-GFP) basal and apical boundaries of the ring, parallel with the growth axis. Measurements of individual ring lengths over a period of 4 h were initiated when separation from the growing tip was discernible. Initiation of measurements of Sep7-GFP intensity at the rings was done identically. To measure the intensity of a given ring, a region of interest was drawn around the boundaries of the ring, and the mean fluorescence value per pixel of the selected area was recorded. An adjacent region of the hypha (lacking a ring) was measured, and this average fluorescence value per pixel was subtracted from the ring area fluorescence value to yield the fluorescence per pixel that was contributed by the ring. The regions of interest were redrawn at every time point to compensate for any reorientation or movement of hyphae or gel pad drift. The variable presence of moving nuclei in the ring's region of interest contributes to noise in these measurements as nuclei have considerably less fluorescence than the cortex or the cytosol. Time-lapse movies were linearly contrast enhanced in Volocity 4.

Visualization of F-Actin, Microtubules, and Chitin

To view F-actin and maintain viability of the Sep7-GFP fluorophore, cells were fixed in a 2% paraformaldehyde (Fisher Scientific, Fair Lawn, NJ) and full media solution at 30°C (unless otherwise indicated). After fixing for 5–10 min, cell cultures were immediately washed with $1\times$ phosphate-buffered saline (PBS) to prevent overfixation and concentrated by centrifugation. Cells were washed again and resuspended in $1 \times$ PBS. To stain for F-actin, cells were incubated with AlexaFluor-Phalloidin568 (Invitrogen) at a concentration of 6.6 μ M and incubated at room temperature in the dark for 1 h. Cells were then washed twice with $1 \times$ PBS and resuspended in ProLong Gold antifade reagent (Invitrogen). To visualize chitin at septa, cells were grown in liquid culture for 14 h, collected via centrifugation, and washed with distilled H2O. Cells were then stained for 5 min with calcofluor white at a final concentration of $167 \mu g/ml$. Cells were washed twice, resuspended in water, and mounted on a glass slide for observation under the microscope.

For cells treated with latrunculin-B (Lat-B; Alexis Biochemicals, San Diego, CA), liquid cultures were grown overnight for \sim 13 h. These cells were incubated for 1 h at 30°C with latrunculin-B at a final concentration of 50 μ M. For visualization of actin and Sep7-GFP, standard procedures were followed.

Treatment of cells with nocodazole (Sigma-Aldrich, St. Louis, MO) involved two separate fixation protocols. To view the Sep7-GFP fluorophore and nuclei response to nocodazole, cells were grown in liquid cultures for \sim 14 h and treated with nocodazole for 2 h at a final concentration of 20 μ g/ml at 30°C. Cells were fixed in a 2% paraformaldehyde and full media solution at 30°C for 15 min. The culture was then concentrated by centrifugation, washed and suspended with $1\times$ PBS, and incubated at room temperature for 15 min with 1:500 Hoechst (1 mg/ml stock; Invitrogen). Stained cells were collected by centrifugation and resuspended in ProLong Gold antifade reagent. To confirm microtubules depolymerized in response to nocodazole, cells were grown and treated with nocodazole as described above. After incubation with nocodazole, cells were fixed for 1 h in formaldehyde (Fisher Scientific) at a final concentration of 3.7%. Cells were then processed for anti-tubulin immunofluorescence as described previously. Cells were incubated overnight at 4°C in a humid chamber with rat-anti- α -tubulin (Serotec, Oxford, United Kingdom) at a 1:50 dilution in $1 \times$ PBS + bovine serum albumin. After washing, cells were incubated 1 h in 2° antibody (1:200 dilution, Alexa Fluor 568-conjugated goat anti-rat antibody) and 1:500 dilution Hoescht (1 mg/ml stock solution). Cells were mounted on slides in ProLong Gold antifade reagent.

RESULTS

Diverse Higher Order Septin Structures Coexist in One Cell

To visualize the septin cortex in living *A*. *gossypii* cells, Sep7p-GFP was expressed from the endogenous locus as the only copy of the gene and imaged by wide-field fluorescence microscopy followed by iterative deconvolution (see *Materials and Methods*). *A*. *gossypii* cells lacking Sep7p do not

assemble septin rings as measured by Cdc11a-mCherry fluorescence (Supplemental Figure 1). Thus, Sep7p is an integral component of the septin cortex in these fungal cells and essential for higher order structures. Sep7-GFP strains have growth rates and morphology identical to the wild type, indicating that this fusion did not interfere with the function of Sep7p (Figure 1R; Helfer and Gladfelter, 2006; data not shown).

A diverse array of higher order septin structures is visible across developing single-celled mycelia (Figure 1, A–Q and R). Near tips of hyphae, Sep7p forms a diffuse cloud in which some faint, thin filaments can be distinguished (Figure 1A). Within \sim 15 μ m of growing tips, a cortical septin ring composed of discrete, thick filaments or "bars" is present in the majority of hyphae (Figure 1, B and C). We call these high order structures rings because they surround the cortex such that in cross section they look like circles (Figure 1K). Similar rings composed of Sep7p bars were observed at intervals ≥ 10 μ m radiating away from the growing tip, toward the center of the cell (Figure 1D). Eighty-six percent of such rings were at this spacing frequency ($\dot{N} = 142$). We call these rings interregion \overline{I} (IR) rings because they are in the middle region of the cell between the growing tip and the spore (Figure 1D). The dimensions of IR rings vary widely within and between cells. Specifically, the ring length, which was measured parallel to the growth axis, was on average 1.88 μ m \pm 0.622 (SD) long (N = 142). Additionally, double rings of short septin bars are found distant from the growing tips at the center of mycelia, which are septation sites (places where cell wall is deposited to generate two cell compartments, Figure 1, E–H).

Lateral branches emerge from the sides of hyphae in proximity to IR septin rings, indicating that one function of the rings may be in promoting branch emergence (Figure 1, I–L; 80% of branches within $<$ 10 μ m of an IR ring; N = 109). Branches have a ring of septin filaments at their base, reminiscent of the septin structures found at the base of dendritic spines in mammalian neurons (Figure 1L; Tada *et al*., 2007; Xie *et al*., 2007). Branch rings have on average smaller dimensions than the IR rings but still are highly variable $(1.44 \pm 0.516$ [SD] μ m; N = 112). Additionally, the filaments of branch bases generally have much lower fluorescence intensity than IR rings. This makes maximum projections of three-dimensional (3D) images of branches seem to have concentrations of septins primarily at the edges and only faint signal across the hypha, but this is because of the

projection. We could reliably count 14–18 bars/ring in branch base rings (Figure 1K) because branches can form perpendicular to the imaging plane, allowing full use of the high X-Y resolution of the microscope ($N = 10$). Often a new IR ring, deposited by the emerging branch tip, is found within 1.0 μ m of these smaller branch rings (Figure 1M).

At lower frequency than the IR and branch base rings, additional distinct higher order structures can be found throughout cells (Figure 1, N–Q). In mature hyphae (20 h after germination) hyphal tips split into two forming a "Yjunction" and two new branches. Here, a variation on the IR ring is seen in which long filaments traverse the bent, "Y" shape extending from the original single axis into one of the two new axes (Figure 1N). Occasionally, an hourglass ring (Figure 1O) is observed that may be an alternative septation structure. In addition, helices seen in vitro with purified mammalian septins are also present here in vivo (Figure 1P) (Kinoshita *et al.*, 2002). Finally, Sep7p can be found in \sim 1- μ m-diameter circles, of similar appearance and dimensions as those observed by Kinoshita and colleagues in mammalian cells when actin is disrupted (Kinoshita *et al*., 2002; Figure 1Q). These may represent a "reserve" pool of spontaneously forming septin structures that self-assemble into circles when they fail to be incorporated into organized higher order rings (see below).

These data demonstrate that morphologically diverse septin rings are found within a single cell (Figure 1R). Clearly, distinct classes of rings can coexist in a cell cortex that is in contact with a common cytoplasm. Many questions emerge about the organization of the septin cortex from these static images. Where are septin rings born? Can they assemble in the middle of hyphae? Do individual rings change dimensions or filament density with time after assembly? Do neighboring rings influence each other's organization? Do all rings persist for similar times? To address these questions, we performed time-lapse microscopy and followed septin rings in growing mycelia over many hours of development.

Septin Rings Assemble at Sites of Growth and Have Constant Dimensions

To measure the behavior of rings through time and space, Sep7-GFP cells were germinated on agar pads and followed for 12–16 h of growth by fluorescence microscopy. The timelapse data demonstrated that all IR rings assemble at growing tips from clouds of diffuse filaments that initially travel

Figure 1. Sep7-GFP localizes to diverse higher order septin structures. Sep7-GFP (AG127)–expressing cells were imaged live after growth overnight on media containing-agarose pad except where noted. All images are maximum projections of image stacks. (A and B) Sep7p at the hyphal tip. (C) A nascent IR ring. (D and E) IR rings. (F–H) Splitting of an IR ring and septation. (I) Sep7p at an incipient branch site. (J) Branch ring. (K) Cross section of branch ring. (L) Branch ring. (M) IR ring formed adjacent to branch ring. (N) IR ring traversing a Y-junction. (O) Hourglass-shaped septin structure at a septum. (P) Septin helix formed under stress. (Q) Cortically associated \sim 1- μ m ectopic ring. (R) Small mycelia in which some examples of the classes of rings shown in A–Q are noted highlighted by letters. (K and P) Cell was fixed with 2% paraformaldehyde. GFP fluorescence (green) and phase contrast outlines (red) are shown. Bars, 5 μ m (A–J and L–O; in A); 2.5 μ m (K, P, and Q; in Q); and 5 μ m (R).

with the growing tip (Supplemental Movies 1 and 2). Faint filaments detach from the tip and coalesce into discrete, resolvable septin filaments. These thick bars of septins are then anchored to the cortex and form a ring that is left in the "wake" of the growing tip (Supplemental Movies 1 and 2). Thus, IR rings never occur in the middle of hyphae but are exclusively assembled at sites of polarized growth. Similarly, side branch rings recruit Sep7 concurrently with the polarized growth generating branch emergence. However, as seen in the still images, branch rings seem to be fainter and shorter than IR rings (Supplemental Movies 1 and 2).

One possible mechanism by which diverse rings could be maintained on the cortex of a common cytoplasm would be if there were a maturation program or "timer" built into the nascent ring. Given the nucleotide binding capacity of septins, such a mechanism could be imagined based on GTP hydrolysis rates of individual subunits. In this case, we

would expect that, upon assembly, there is a consistent time period from which a ring transitions from its initial appearance into different morphologies. However, splitting and disassembly of rings that were observed from their birth were rare events in the time-lapse movies of young, actively growing cells. We only observed splitting for six of 300 rings that were tracked from their assembly, so it was not possible to obtain significant measurements on the transition timing. Within this small population, no consistent timing pattern for transitions could be observed. We could see that when a cell showed signs of stress such as the appearance of vacuoles, there was a nearly instantaneous transition of many IR rings $(2 h old)$ to split double rings as well as the formation of \sim 1- μ m circles (Supplemental Movie 3). This indicates that rings of very different ages all have a similar potential to transition and change their higher order structures. These data suggest that an intrinsic timer that is based solely on the

Figure 2. Septin rings do not change dimensions with time. Single *SEP7-GFP* (AG127) cells were observed using 3D time-lapse fluorescence microscopy. Image stacks were acquired every 20 min. All images were acquired and processed identically. The sequences of images above the graphs are examples of an IR ring (A) and a branch ring (B) that were measured. (A) Individual IR septin rings were measured for 4 h after their establishment. $N = 50$. (B) Branch rings were measured for 4 h after their establishment. $N = 30$. The small fluctuations in these measurements reflect minor shifts in the orientation of the hypha containing the ring as it grows. GFP fluorescence (green) and phase contrast cell outlines (red) are shown. Bars, 5 μ m.

age of the septin complex is unlikely to be the primary basis for the diversity in septin ring organizations.

The time-lapse approach also enabled us to analyze the basis for variability in dimensions of the septin rings within different classes. The variability seen in the population could arise from heterogeneity in the initial dimensions of the ring at assembly or dynamics during maturation that led to the contraction or expansion (potentially polymerization or depolymerization) of filaments within the ring. Our data demonstrate that the initial dimensions of rings at their inception were maintained through the ring's lifetime until either disassembly or splitting (Figure 2, A and B). Individual rings did not change dimensions with time except on the rare occasion when a ring splits into two rings at septation sites (Figure 1, G and H, and Supplemental Movies 3 and 4). Thus, at birth, an individual ring's maximum size is established, however, the absolute size varies across rings. The fixed dimensions are indicative of a septin "cap," potentially analogous to capping proteins for actin, that constrains the expansion of the cortex after assembly.

Septin IR Rings Assimilate New Septin Subunits over Extended Time Periods

The constant dimensions of septin rings after assembly suggest that factors at the growing tip promote assembly and then rings are "static" until unknown cues trigger septation in a small subset of rings. Alternatively, rings may be constant in geometry but change in either the composition or dynamics of individual septin subunits and thus mature through time. These alternative models for maturation prompted us to measure the rate of incorporation of septin protein through the life of a ring.

To assay Sep7p incorporation, the fluorescence intensity of Sep7-GFP at individual septin rings was determined for IR rings through time. We calculated an average fluorescence intensity from a three-dimensional reconstructed image at each time point (initial ring appearance to 240 min old). To control for machine noise and fluorescence in the cytosol underlying the septin cortex, a background fluorescence intensity from neighboring cytosol was measured at each time point and subtracted from the average fluorescence of the ring (see *Materials and Methods*). To measure the change in fluorescence of IR rings through time, the initial normalized fluorescence value for each ring was subtracted from each time point in the series. IR rings on average increase 2.5 times in their fluorescence intensity over 4 h; however, this increase varied from as little as 0.5 times to >7.0 times the starting intensity (Figure 3, A–C; N = 50 rings). The rate of increase in fluorescence we observed was significant based on a Student's t test ($p < 0.001$) in com-

Figure 3. Sep7-GFP intensity increases with time at IR septin rings but not at branch rings. Single *SEP7-GFP* (AG127) cells were observed using 3D time-lapse fluorescence microscopy. Image stacks were acquired every 20 min and were processed identically. (A and E) Images show examples of an IR ring (A) and a branch ring (E) that were measured. The GFP fluorescence (green) and phase contrast cell outlines (red) are shown. Bars, 5 μ m. (B and F) Cumulative change in fluorescence intensity of Sep7-GFP at IR rings (B; N = 50) and branch rings (F; $N = 30$) was measured for 4 h after establishment. Change in fluorescence intensity was calculated by subtracting the intensity of the ring immediately after establishment (t = 0) from the intensity measured at a given time point. Intensity values were corrected for background fluorescence of the cytosol. Values are given in arbitrary fluorescence units (au). The black lines represent the median cumulative change for

parison with a null hypothesis of no increase in fluorescence. The rates of change in fluorescence vary significantly between rings ($p < 0.01$) but are nonetheless linear for all rings measured regardless of differences in their -fold change in fluorescence intensity (Figure 3, B and C). The linear behavior of the rate of fluorescence change in IR rings suggests that the incorporation of new subunits is likely due to filament polymerization or the stoichiometric association of additional free subunits with the existing polymers.

For an individual ring, the rates of intensity change are constant through time, indicating that addition of Sep7p to rings does not reach saturation in this interval. This suggests that subunit addition may be mechanistically similar both in very young rings and mature rings. Additionally, there is no correlation between the rate of fluorescence intensity change and the initial fluorescence intensity (Figure 3D; $\dot{R} = 0.20$, $N = 50$). This indicates that the variation in rate of change between rings is not arising due to different starting amounts of Sep7p. Furthermore, the rate of fluorescence incorporation for a given ring does not correlate with its length ($R = 0.11$, $N = 50$) or due to its proximity to another septin ring ($R = 0.19$, $N = 20$). Thus, the rate of addition of new subunits is not determined by the amount of Sep7p in a preexisting ring or the length of that ring.

In striking contrast to IR rings, when similar intensity measurements were made on the rings at the bases of branches (such as the type in Figure 1, I–L) Sep7p intensity did not increase and thus had fluorescence change close to or at zero (Figure 3, E–G; $N = 30$ rings). The median rate of change in this class of rings is slightly negative, potentially because of photobleaching over time or loss of subunits. Branch rings fail to increase in intensity even when nearby $(<$ 10 μ m away) IR rings do increase fluorescence intensity, indicating that this lack of incorporation is not likely because of local limitation in the concentration of free subunits.

Variable Septin Morphology Is Not Because of the Presence or Absence of Different Septin Subunits within a Ring

Given the variability in intensity and ring organization that we observed in one cell, we hypothesized that different septin structures within *A*. *gossypii* could arise because of the presence or absence of one or more of the six septin proteins in different classes of rings. This idea is attractive because of the ability of only two or three septin subunits to polymerize in vitro and assemble higher order structures in other species. To test this hypothesis, we tagged each individual septin subunit with a fluorescent protein and determined whether each septin subunit was found in all classes of septin rings. Each of the septins examined (Cdc3, Cdc10, Cdc11a, Cdc12, and Sep7) was found in all classes of rings,

Figure 4. All classes of septin rings contain all septin subunits. *A*. *gossypii* strains expressing Cdc11a-mCherry from the endogenous locus and either Sep7-GFP (AG232), Cdc10-YFP (AG306), or Cdc12- Venus(AG308) expressed using their endogenous promoter from a replicating plasmid were imaged in live cells grown overnight on agarose media pads. mCherry fluorescence (red), GFP fluorescence (green), Venus fluorescence (yellow), and phase contrast outlines (blue) are shown. Bar, $5 \mu m$.

uniformly across the entire mycelia (Figure 4 and Supplemental Figure 2). To address whether two septin subunits precisely colocalized based on fluorescence signals, we coexpressed pairs of septin subunits that were tagged with different fluorescent proteins.

Representative colocalizations for Sep7-GFP and Cdc11 mCherry, Cdc10-YFP and Cdc11-mCherry, and Cdc12-YFP and Cdc11-mCherry are shown in Figure 4. In all pairwise combinations tested, the septins exactly colocalize in rings. Thus, as detectable by fluorescence, individual septin subunits are symmetrically and uniformly distributed in all classes of septin rings. These data indicate that all expressed septins are incorporated into all rings and that variable higher order organizations do not arise because of the incorporation or exclusion of specific septin subunits.

Septin Rings Are Maintained Independent of Actin or Microtubule Cytoskeletons

In mammalian cells, higher order septin organization has been linked to both the actin and microtubule cytoskeletons. We therefore tested the hypothesis that variability in ring structure arises because of associations with the actin or microtubule cytoskeletons. First, we observed the colocalization of F-actin and Sep7-GFP in normal growth conditions. The faint septin filaments form at growing tips where actin cables are nucleated; however, we could not resolve individual septin filaments juxtaposed to actin cables. New IR rings are located just behind the concentration of F-actin patches that congregate at growing tips (Figure 5A). Rings of F-actin were found at only a subset of IR and branch base rings. These F-actin rings are located at septin IR rings in the center of cells closest to the spores that are the most mature.

Figure 3 (cont). the population. Fifty percent of the measured rings are in the dark gray regions, and 80% of the rings fall in the combined light gray and dark gray regions. (C and G) The box and whisker plots show the distribution of slopes of the linear regression of fluorescence change at individual rings. The black line denotes the median value and the asterisk shows the mean value. The box is the middle 50% of the data, and the whiskers extend to the most extreme data point, no more than 1.5 times the interquartile range. Rates are given in arbitrary fluorescence units per minute (au/min). (D and H) The slopes of the linear regression of fluorescence changes for a ring are graphed against the initial fluorescence at that ring for each individual septin ring analyzed. Values are given in arbitrary fluorescence units and rates are given in arbitrary fluorescence units per minute. In both cases, R values are insignificant.

Figure 5. F-Actin rings are asymmetrically localized at septin rings. (A) Colocalization of Sep7-GFP and F-actin. *SEP7-GFP* cells (AG127) were grown overnight in liquid media at 30°C. Cells were fixed with 2% paraformaldehyde and the F-actin was visualized using AlexaFluor-Phalloidin568. (B) Actin rings localized to the basal end (top) or center (bottom) of established septin rings. Arrows point to the hyphal tip and branch point. GFP fluorescence (green), Alexa568-Phalloidin (red), and phase contrast cell outlines (blue) are shown. Bar is 3μ m. (C) Actin ring position was scored relative to the associated septin ring (N = 141).

Remarkably, the actin rings are not localized in the center of septin rings but instead were on the basal end (side closest to the center of the cell, away from the growing tip) in 67% of rings ($N = 141$). In the remaining cases, actin rings are in the center of the septin ring but never on the apical side of the septin ring. The asymmetric association related to the

length of the ring such that rings with an actin ring at the basal side are on average 0.90 \pm 0.19 (SD) μ m long, whereas septin rings with actin rings in the middle are on average $0.51 \pm 0.13 \mu m$ (N = 82; the smaller overall dimensions compared with the live imaging is because of fixation-induced cell shrinkage). These data indicate that there is some

Figure 6. F-actin is required for septin ring assembly but not maintenance. *SEP7-GFP* cells (AG127) were grown overnight in liquid AFM at 30°C. (A) Before heat shock, cells were fixed with 2% paraformaldehyde, and the F-actin was visualized using AlexaFluor-Phalloidin568. (B) Cells were transferred to 42°C for 20 min to depolarize the actin cytoskeleton and then processed for Factin visualization. (C) Heat-shocked cells were returned to 30°C to recover for 20 min and then processed for F-actin visualization. (D) Cells were incubated for 1 h in growth media with latrunculin-B (final concentration, 50 μ M) and then processed for F-actin visualization. GFP fluorescence (green), AlexaFluor-Phalloidin568 fluorescence (red), and phase contrast cell outlines (blue) are shown. Bar, 5 μ m.

polarity within a ring as to the ability to recruit F-actin and that longer rings tend to be more asymmetric (Figure 5). Given that actin rings are observed only in mature septin rings that may already be many hours old, this septin ring asymmetry must be maintained for an extended time or established well after the initial assembly of the ring.

We perturbed the normal organization of F-actin with heat shock by transferring cells to 42°C for 20 min. This treatment leads to a rapid depolarization of the actin cytoskeleton in which cables and rings disassembled and Factin patches redistributed uniformly along hyphae (Figure 6, A and B). On heat shock, Sep7-GFP fails to accumulate at growing tips (Figure 6B). All other higher order septin structures persist and are found with similar frequency to normal conditions (with the exception of an increase in the frequency of \sim 1 μ m circles; Figure 6B). When returned to 30°C, tip-associated septins accumulate when cell polarity is reestablished, usually within 20 min of recovery (Figure 6C). This indicates that only the initial tip-associated assembly of septin filaments requires polarized F-actin.

To determine whether the loss of septin filaments at tips is directly due to the loss of actin polarity or due to a more

general response to temperature stress, we incubated cells in 50 μ M Lat-B, which is sufficient to eliminate all F-actin in both cables and patches in cells (Figure 6D). Similar to heat shock, septin filaments are not assembled at tips in the presence of Lat-B, but all other classes of assembled rings persist throughout the cell (Figure 6D).

Additionally, during the incubations both at high temperature and in Lat-B, we observed a profusion of \sim 1- μ m septin circles. This suggests that these structures do not require actin polarity or cables for assembly. On return to 30°C after heat shock, the tip-associated septin filaments reestablish to all growing tips, concomitant with the disappearance of the small septin circles. This observation led us to speculate that these circles may form spontaneously at a sufficient concentration of free septins in the cytosol. In support of this idea, we have seen these circles in live cells when tip growth and IR ring assembly stop transiently. We have also observed them disappear just before a new branch ring forming (Supplemental Movies 4 and 5). We speculate that the \sim 1- μ m circles are likely self-assembling structures that form above a certain cytosolic concentration of septin oligomers or polymers. Thus, the majority of septin struc-

Figure 7. Microtubules are not required for septin ring assembly or maintenance. Cells expressing Sep7-GFP (AG127) were grown overnight in liquid AFM media at 30 \degree C (t = 0) and then were treated with 20 μ g/ml nocodazole for 120 min and fixed for visualization of Sep7-GFP (A) or anti- α -tubulin immunofluorescence (B) in combination with Hoechst staining of DNA. Sep7-GFP (green), anti- α -tubulin (red), and Hoechst (blue) stainings are shown. Bars, 5 μ m.

tures are maintained in the absence of polarized F-actin; however, assembly of nascent rings requires cell polarity.

To determine whether microtubule dynamics or polymerization are required for septin organization, we incubated Sep7-GFP–expressing cells in the microtubule-destabilizing drug nocodazole. Under normal growth conditions, *A*. *gossypii* cells have intranuclear spindle microtubules, short cytosolic astral microtubules that emanate from the spindle pole body, and extensive cytosolic microtubules on which nuclei are transported for long-range migration toward the growing tip (Alberti-Segui *et al*., 2001; Gladfelter *et al*., 2006). Incubation in nocadazole leads to disassembly of all microtubules and induces an arrest of nuclei at G2/M in the division cycle, although polarized growth of hyphal tips continues at normal rates (Figure 7A; Gladfelter *et al*., 2006). On loss of microtubules, all higher order septin structures are retained, and new septin filaments are still visible at growing tips (Figure 7B). These data indicate that neither the progression of nuclei through the division cycle nor the microtubule cytoskeleton is required for any detectable feature of septin assembly or organization. This separation between septin organization and cell cycle progression is similar to what is observed in differentiated mammalian cells such as neurons where septins assemble and mature in G_0 .

Specific Ring Morphologies Require the Activity of Different Regulatory Proteins

The variable septin organization observed in *A*. *gossypii* is not because of an intrinsic timing program built into the septin ring, the presence or absence of specific septin subunits, or the actin or microtubule cytoskeletons. We hypothesized that distinct septin structures form at the cortex of a common cytoplasm because of the localized activity of septin regulators. Regulators such as kinases could deliver posttranslational modifications to septins that either lead to the assembly of different classes of rings or different maturation properties of individual rings. Ring-autonomous maturation steps that may be regulated include the initial anchoring to the cortex, the addition of new septin subunits to a preexisting ring or the compression of filaments at a splitting ring. We predicted that mutation of such a regulator may disrupt only a subset of rings that require that activity for

their genesis or maintenance, whereas the remaining classes are unaffected.

We tested this hypothesis by deleting the septin-associated kinases *ELM1* and *GIN4* and a septin-interacting factor encoded by *NAP1* in cells expressing Sep7p-GFP. Notably, deletion of either *GIN4* or *ELM1* leads to complete loss of all IR rings found along main hyphae ($N \ge 500$ hypha examined for each mutant). However, in both mutants there are still detectable fine filaments at tips, \sim 1- μ m circles, and branch base rings (Figure 8). Mutants lacking Elm1p have more intense tip associated filaments, whereas $\sin 4\Delta$ mutants have tip filaments comparable with wild type, indicating that potentially these kinases play different roles in the establishment of IR rings. In both mutants, the branch base rings had a similar spectrum of dimensions and fluorescence intensities as observed for wild-type cells, suggesting that these regulators are not required for any feature of this class of rings. We confirmed that these phenotypes were because of altered organization of the entire septin ring and did not simply delocalize Sep7p by observing the same phenotypes in mutants expressing a Cdc11a-mCherry (Supplemental Figure 3). Thus, these kinases are dispensable for the formation of branch base rings and filament assembly at tips but are required for IR ring morphogenesis.

Cells lacking these kinases showed several morphological defects in addition to loss of the septin IR rings. *elm1* mutant hyphae (3.04 \pm 0.03 [SEM] μ m; N = 50) and *gin4* Δ mutant hyphae (3.92 \pm 0.04 μ m; N = 55) are both significantly more narrow compared with wild-type cells $(4.17 \pm$ $0.05 \mu m$; N = 66; p < 0.001 based Student's t test). Both $elm1\Delta$ and $gin4\Delta$ mutants lack any septation sites as monitored by calcofluor staining of the cell wall (Supplemental Figure 4). In response to localized cell stress or death, septa in wild-type cells act as "bulk-heads" and wall off the damaged cell from the rest of the mycelia (Supplemental Movie 3 and Supplemental Figure 5). In contrast, in the $\sin 4\Delta$ and $elm1\Delta$ mutants, the cell stress phenotype spreads across the entire mycelia and is not locally contained (Supplemental Figure 5). Finally, both mutants showed diminished colony radial growth rates. $\sin 4\Delta$ is 7.55 \pm 0.22 (SEM) mm/d, *elm*1 Δ is 7.39 \pm 0.36 mm/d compared with wild type, which is 9.75 ± 0.31 mm/d as measured for four independent colonies over 9 d. These growth differences may be because of both the hyphal morphology defects and the inability to compartmentalize hypha with septa because of the lack of IR rings.

To determine whether Elm1p and Gin4p function in the assembly or maintenance of IR rings, we followed septin organization in *elm1* Δ and *gin4* Δ cells through time (Supplemental Movies 6 and 7). We observed that IR rings do not assemble in either mutant. Septins concentrate at the tips in diffuse filaments in both mutants. These filaments detach from the growing tip but only transiently coalesce into faint, thin bars that are associated with the cortex. In no case are normal IR rings observed at a concentrated zone behind the growing tip. In contrast, branch ring assembly and maintenance does not differ from wild-type cells. Thus, the assembly of rings fails in both of these mutants, with a clear defect in anchoring and construction of the higher order bars in IR rings.

Deletion of *NAP1* produces highly misorganized and asymmetric IR rings but, like the mutants described above, has no effect on the formation or organization of branch base rings. This suggests that Nap1p is important for the symmetric and constant dimensions that are imparted to a ring at assembly. Alternatively, symmetric rings may form transiently and then disassemble asymmetrically in the

Figure 8. Deletion of septin regulators disrupts IR rings but not branch rings. Live cell fluorescence images of wild-type *SEP7-GFP* (AG127), *gin4 SEP7-GFP* (AG121), *elm1 SEP7-GFP* (AG120), and *nap1 SEP7-GFP* (AG209) cells grown overnight on agarose pads. Asterisks, squares, and arrows highlight the presence of tip-associated filaments, interregion rings, and branch rings, respectively. In the *nap1* a panel, an asymmetric interregion ring is highlighted and enlarged. GFP fluorescence (green) and phase contrast cell outlines (red) are shown. Bar, 5 μ m.

absence of Nap1p. Time-lapse analysis of Sep7-GFP in the $nap1\Delta$ mutant strains reveals that rings initially assemble with asymmetric dimensions, indicating a role for Nap1p in establishing IR ring dimensions and regular geometry (Figure 9).

Figure 9. *nap1*∆ mutants assemble asymmetric IR rings. Images from a time-lapse series in which *nap1* (AG314) cells were grown on an agar gel pad and imaged every 30 min. Sep7-GFP is asymmetrically deposited in IR rings as they form (arrow and arrowhead). Bar, 5 μ m.

DISCUSSION

With the data presented here, we have established a new model system for the study of higher order septin structures. This fungal system complements current models for septin analysis in several important ways. Diverse, persistent, and distinct higher order structures form in the context of branching and polarized cell morphology (Tada *et al*., 2007; Xie *et al*., 2007). The coexistence and persistence of multiple, distinct septin structures in these branched fungal cells are remarkably similar to that of septins observed in the dendritic spines of mammalian neurons. We show that the long-lived rings are not static but rather change in their appearance and incorporate new subunits over many hours of existence. Importantly, these changes occur independently of the cell cycle. Thus, this is a powerful, genetically tractable model system in which to study mechanisms of in vivo assembly and maturation of the septin cortex without complication of cell cycle remodeling cues. Additionally, we show that IR and branch rings differ in their appearance, maturation and regulatory requirements. This indicates that spatially restricted signaling can produce functionally and morphologically distinct septin complexes within a single cell.

Septin complexes in *A*. *gossypii* initially assemble into fine filaments at growing hyphal tips and travel with the growth zone. After growing with the tip for $10-15 \mu$ m, an Elm1p/ Gin4p kinase-dependent signal triggers the coalescence of filaments into thicker bundles and anchoring of complexes to the cortex to form stable IR rings. The role of these regulators may be to bundle filaments into higher order bars, to detach filaments from the site of polarized growth and/or to anchor septins to the plasma membrane. Interestingly, in *Candida albicans* hyphae, septins also initially congregate in a diffuse cloud at the growing hyphal tip (Sudbery, 2001; Gonzalez-Novo *et al*., 2004; Wightman *et al*., 2004; Gonzalez-Novo *et al*., 2008). Coordinately with the cell cycle, the septin clouds then coalesce into double rings to form septa that generate uninucleate compartments in the hyphae. Interestingly, Gin4p in *C*. *albicans* is dispensable for the tip-associated septin assembly and also the most basal septin structure formed at the germ bubble (Wightman *et al*., 2004). In *C*. *albicans* the dissociation of the cloud from the tip and formation of the septa is dependent upon Gin4p kinase, similar to what we have observed for *A*. *gossypii*, suggesting that the membrane anchoring event may be conserved among fungi and potentially animals. The abrupt kinasedependent transition from faint, cytosolic filaments to bundles of thick bars fixed to the cortex, will enable future experiments to probe the molecular basis for septin/membrane connections. This is especially important because the mechanisms directing septin association to membranes are not known in any system, and we now have specific mutations that block this process.

The newly assembled IR rings vary in their initial size, intensity, and capacity to add subunits with time. However, each of these physical features function independently of one another because there is no correlation between any of these traits across populations of rings. Septin rings composed of bars similar to those in the IR rings are also seen in *S*. *cerevisiae* shmoos and *gin4* mutants, and in *C*. *albicans* at the base of hyphal cells and in chlamydospores (Giot and Konopka, 1997; Sudbery, 2001; Wightman *et al*., 2004; Martin *et al*., 2005). However, none of these bar-structures require Gin4p kinase for assembly and/or are as persistent as the bars in IR rings that we observe. Therefore, we predict that there may be a different regulatory basis for the organization of septins into bars in the IR rings of *A*. *gossypii*. The septin bars we observe are spaced regularly around the cortex in a ring, and the number of bars per septin ring seems constant over an area, raising the question of how such uniform spacing is achieved. Potentially, there may be uncharacterized septin associated proteins that function similarly to microtubule-associated proteins and act as spacers to delineate the zones between septin bars.

After assembly and anchorage, the IR rings persist for indefinite periods (over multiple ≥ 16 -h time series, only 6/300 rings disappeared). Notably, these IR rings continue to incorporate new septin subunits; in many cases rings more than double in their fluorescence intensity over 4 h of observation. This kind of maturation of the septin cortex has not been noted in other model fungi but may be a trait in higher eukaryotes where septin structures persist for extended periods. We predict that this additional protein incorporation could represent new filament polymerization or nonpolymerized Sep7p associating peripherally with septin filaments already organized in the ring. New filaments may polymerize in situ at a preassembled septin cortex or cytosolic filaments may intercalate into the preexisting higher order structure. It is also possible that the Sep7p subunits

associate with the assembled structures without actually incorporating into filaments.

It is intriguing that new protein is added exclusively at bars and not in the spaces between the bars, suggesting that the preexisting structure is a template for the addition/ polymerization of new protein. The rate of subunit addition varies but is always linear, indicating that the template/ nucleator recruits a stoichiometric amount of either free subunit or polymer. Future experiments using photoactivatable GFP fused to septins and fluorescence recovery after photobleaching of the septin cortex will be useful in monitoring the position in the ring where addition takes place. Furthermore, there is no elongation of a septin bar with time suggesting that there maybe a capping protein prohibiting expansion from the ends of the existing polymer. The irregular alignment and variable lengths of filaments in $nap1\Delta$ mutants suggest that Nap1p may be required for capping the ends of the rings.

A notable trait of maturing IR rings is that they recruit F-actin rings asymmetrically and preferentially position actin at the basal (opposite from growth tip) end of the septin bars. This observation is striking given the known nonpolar nature of individual septin filaments formed in vitro (John *et al*., 2007; Sirajuddin *et al*., 2007; Bertin *et al*., 2008). Interestingly, in *A*. *nidulans*, the septin ring may disassemble in an asymmetric manner, with the basal side disappearing and the apical side persisting after septation (Westfall and Momany, 2002). In *S*. *cerevisiae*, dozens of proteins are asymmetrically localized to only one side of the septin collar (Gladfelter *et al*., 2001). In no case is the basis of the asymmetry known, although data in *S*. *cerevisiae* suggest that asymmetry is not due to the septins themselves but must be layered upon the septins by additional proteins (Vrabioiu and Mitchison, 2007). Our data further support that functional asymmetry can arise from a higher order septin structure in which septins are presumably symmetrically organized and suggest that this property is evolutionarily conserved.

Branch rings assemble near IR rings, indicating that the IR rings may serve as landmarks for the location of branch emergence. Cells lacking all septin structures or missing only IR rings still form branches but of aberrant morphology and positioning (Supplemental Figure 1 and Figures 8 and 9; Helfer and Gladfelter, 2006). Remarkably, the branch base septin rings do not add subunits with time and do not require the Elm1/Gin4 kinase for their assembly or maintenance. Therefore, the branch rings are morphologically distinct from IR rings, yet they generally form less than a few microns away from IR rings at the cortex of a common cytoplasm. New branch rings do not "poach" septin subunits from their neighboring IR rings, but in fact, neighboring IR rings will continue to increase in fluorescence intensity even while side branch rings form nearby. Presumably the different rings have access to common cytoplasmic pools of both septin subunits and regulatory proteins. This raises the question of how these different classes of septin rings are structurally and functionally distinguished despite their residence in the same space.

These data represent the debut of a new model system for the analysis of in vivo septin assembly and organization. We have demonstrated that the septin cortex shows great plasticity within a single cell. *A*. *gossypii* septin assembly, maturation, and organization do not require the nuclear cycle so this will be an important system for studying cell cycleindependent regulation of septins. The different septin structures and behaviors combined with the battery of mutants that we have developed make *A*. *gossypii* a great resource for biochemical and quantitative microscopy approaches to septin function in the future. We are well poised to understand how septin proteins may be modified to attach to membranes and to form diverse and persistent higher order structures in cells.

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REFERENCES

Alberti-Segui, C., Dietrich, F., Altmann-Johl, R., Hoepfner, D., and Philippsen, P. (2001). Cytoplasmic dynein is required to oppose the force that moves nuclei towards the hyphal tip in the filamentous ascomycete *Ashbya gossypii*. J. Cell Sci. *114*, 975–986.

Ayad-Durieux, Y., Knechtle, P., Goff, S., Dietrich, F., and Philippsen, P. (2000). A PAK-like protein kinase is required for maturation of young hyphae and septation in the filamentous ascomycete *Ashbya gossypii*. J. Cell Sci. *113*, 4563–4575.

Barral, Y., Mermall, V., Mooseker, M. S., and Snyder, M. (2000). Compartmentalization of the cell cortex by septins is required for maintenance of cell polarity in yeast. Mol. Cell *5*, 841–851.

Baudin, A., Ozier-Kalogeropoulos, O., Denouel, A., Lacroute, F., and Cullin, C. (1993). A simple and efficient method for direct gene deletion in *Saccharomyces cerevisiae*. Nucleic Acids Res. *21*, 3329–3330.

Bertin, A., McMurray, M. A., Grob, P., Park, S. S., Garcia, G., 3rd, Patanwala, I., Ng, H. L., Alber, T., Thorner, J., and Nogales, E. (2008). *Saccharomyces cerevisiae* septins: supramolecular organization of heterooligomers and the mechanism of filament assembly. Proc. Natl. Acad. Sci. USA *105*, 8274–8279.

Caviston, J. P., Longtine, M., Pringle, J. R., and Bi, E. (2003). The role of Cdc42p GTPase-activating proteins in assembly of the septin ring in yeast. Mol. Biol. Cell *14*, 4051–4066.

Dietrich, F. S., *et al*. (2004). The *Ashbya gossypii* genome as a tool for mapping the ancient *Saccharomyces cerevisiae* genome. Science *304*, 304–307.

Dobbelaere, J., Gentry, M. S., Hallberg, R. L., and Barral, Y. (2003). Phosphorylation-dependent regulation of septin dynamics during the cell cycle. Dev. Cell *4*, 345–357.

Egelhofer, T. A., Villen, J., McCusker, D., Gygi, S. P., and Kellogg, D. R. (2008). The septins function in G1 pathways that influence the pattern of cell growth in budding yeast. PLoS ONE *3*, e2022.

Finger, F. P., Kopish, K. R., and White, J. G. (2003). A role for septins in cellular and axonal migration in *C*. *elegans*. Dev. Biol. *261*, 220–234.

Frazier, J. A., Wong, M. L., Longtine, M. S., Pringle, J. R., Mann, M., Mitchison, T. J., and Field, C. (1998). Polymerization of purified yeast septins: evidence that organized filament arrays may not be required for septin function. J. Cell Biol. *143*, 737–749.

Giot, L., and Konopka, J. B. (1997). Functional analysis of the interaction between Afr1p and the Cdc12p septin, two proteins involved in pheromoneinduced morphogenesis. Mol. Biol. Cell *8*, 987–998.

Gladfelter, A. S. (2006a). Control of filamentous fungal cell shape by septins and formins. Nat. Rev. Microbiol. *4*, 223–229.

Gladfelter, A. S. (2006b). Nuclear anarchy: mitosis in multinucleated cells. Curr. Opin. Microbiol. *9*, 547–552.

Gladfelter, A. S., Hungerbuehler, A. K., and Philippsen, P. (2006). Asynchronous nuclear division cycles in multinucleated cells. J. Cell Biol. *172*, 347–362.

Gladfelter, A. S., Pringle, J. R., and Lew, D. J. (2001). The septin cortex at the yeast mother-bud neck. Curr. Opin. Microbiol. *4*, 681–689.

Gonzalez-Novo, A., Correa-Bordes, J., Labrador, L., Sanchez, M., Vazquez de Aldana, C. R., and Jimenez, J. (2008). Sep7 is essential to modify septin ring dynamics and inhibit cell separation during *Candida albicans* hyphal growth. Mol. Biol. Cell *19*, 1509–1518.

Gonzalez-Novo, A., Jimenez, J., Garcia, M. J., Rios-Serrano, I., Pla, J., Jimenez, A., and Sanchez-Perez, M. (2004). Dynamics of CaCdc10, a septin of *Candida albicans*, in living cells and during infection. Int. Microbiol. *7*, 105–112.

Haarer, B. K., and Pringle, J. R. (1987). Immunofluorescence localization of the *Saccharomyces cerevisiae CDC12* gene product to the vicinity of the 10-nm filaments in the mother-bud neck. Mol. Cell. Biol. *7*, 3678–3687.

Hall, P. A., and Russell, S. E. (2004). The pathobiology of the septin gene family. J. Pathol. *204*, 489–505.

Hall, P. A., Todd, C. B., Hyland, P. L., McDade, S. S., Grabsch, H., Dattani, M., Hillan, K. J., and Russell, S. E. (2005). The septin-binding protein anillin is overexpressed in diverse human tumors. Clin. Cancer Res. *11*, 6780–6786.

Hartwell, L. H. (1971). Genetic control of the cell division cycle in yeast. IV. Genes controlling bud emergence and cytokinesis. Exp. Cell Res. *69*, 265–276.

Helfer, H., and Gladfelter, A. S. (2006). AgSwe1p regulates mitosis in response to morphogenesis and nutrients in multinucleated *Ashbya gossypii* cells. Mol. Biol. Cell *17*, 4494–4512.

Ihara, M., Tomimoto, H., Kitayama, H., Morioka, Y., Akiguchi, I., Shibasaki, H., Noda, M., and Kinoshita, M. (2003). Association of the cytoskeletal GTPbinding protein Sept4/H5 with cytoplasmic inclusions found in Parkinson's disease and other synucleinopathies. J. Biol. Chem. *278*, 24095–24102.

Ihara, M., *et al*. (2007). Sept4, a component of presynaptic scaffold and Lewy bodies, is required for the suppression of alpha-synuclein neurotoxicity. Neuron *53*, 519–533.

John, C. M., *et al*. (2007). The *Caenorhabditis elegans* septin complex is nonpolar. EMBO J. *26*, 3296–3307.

Johnson, E. S., and Blobel, G. (1999). Cell cycle-regulated attachment of the ubiquitin-related protein SUMO to the yeast septins. J. Cell Biol. *147*, 981–994.

Kaufmann, A. A plasmid collection for PCR-based gene targeting in the filamentous ascomycete *Ashbya gossypii*. Fung. Genet. Biol. (In Press).

Kinoshita, M., Field, C. M., Coughlin, M. L., Straight, A. F., and Mitchison, T. J. (2002). Self- and actin-templated assembly of mammalian septins. Dev. Cell *3*, 791–802.

Martin, S. W., Douglas, L. M., and Konopka, J. B. (2005). Cell cycle dynamics and quorum sensing in *Candida albicans* chlamydospores are distinct from budding and hyphal growth. Eukaryot. Cell *4*, 1191–1202.

McElver, J., and Weber, S. (1992). Flag N-terminal epitope overexpression of bacterial alkaline phosphatase and Flag C-terminal epitope tagging by PCR one-step targeted integration. Yeast *8* (special issue), S627.

McMurray, M. A., and Thorner, J. (2008). Septin stability and recycling during dynamic structural transitions in cell division and development. Curr. Biol. *18*, 1203–1208.

Mortensen, E. M., McDonald, H., Yates, J., 3rd, and Kellogg, D. R. (2002). Cell cycle-dependent assembly of a Gin4-septin complex. Mol. Biol. Cell *13*, 2091– 2105.

Pan, F., Malmberg, R. L., and Momany, M. (2007). Analysis of septins across kingdoms reveals orthology and new motifs. BMC Evol. Biol. *7*, 103.

Rodal, A. A., Kozubowski, L., Goode, B. L., Drubin, D. G., and Hartwig, J. H. (2005). Actin and septin ultrastructures at the budding yeast cell cortex. Mol. Biol. Cell *16*, 372–384.

Russell, S. E., and Hall, P. A. (2005). Do septins have a role in cancer? Br. J. Cancer *93*, 499–503.

Sambrook, J., and Russell, D. W. (2001). Molecular Cloning: A Laboratory Manual, Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press.

Schmitz, H. P., Kaufmann, A., Köhli, M., Laissue, P. P., and Philippsen, P. (2006). From function to shape: a novel role of a formin in morphogenesis of the fungus *Ashbaya gossypii*. Mol. Biol. Cell *17*, 130–145.

Sirajuddin, M., Farkasovsky, M., Hauer, F., Kuhlmann, D., Macara, I. G., Weyand, M., Stark, H., and Wittinghofer, A. (2007). Structural insight into filament formation by mammalian septins. Nature *449*, 311–315.

Sitz, J. H., Baumgartel, K., Hammerle, B., Papadopoulos, C., Hekerman, P., Tejedor, F. J., Becker, W., and Lutz, B. (2008). The Down syndrome candidate dual-specificity tyrosine phosphorylation-regulated kinase 1A phosphorylates the neurodegeneration-related septin 4. Neuroscience.

Sudbery, P. E. (2001). The germ tubes of Candida albicans hyphae and pseudohyphae show different patterns of septin ring localization. Mol. Microbiol. *41*, 19–31.

Tada, T., Simonetta, A., Batterton, M., Kinoshita, M., Edbauer, D., and Sheng, M. (2007). Role of Septin cytoskeleton in spine morphogenesis and dendrite development in neurons. Curr. Biol. *17*, 1752–1758.

Takizawa, P. A., DeRisi, J. L., Wilhelm, J. E., and Vale, R. D. (2000). Plasma membrane compartmentalization in yeast by messenger RNA transport and a septin diffusion barrier. Science *290*, 341–344.

Versele, M., Gullbrand, B., Shulewitz, M. J., Cid, V. J., Bahmanyar, S., Chen, R. E., Barth, P., Alber, T., and Thorner, J. (2004). Protein-protein interactions governing septin heteropentamer assembly and septin filament organization in *Saccharomyces cerevisiae*. Mol. Biol. Cell *15*, 4568–4583.

Versele, M., and Thorner, J. (2004). Septin collar formation in budding yeast requires GTP binding and direct phosphorylation by the PAK, Cla4. J. Cell Biol. *164*, 701–715.

Versele, M., and Thorner, J. (2005). Some assembly required: yeast septins provide the instruction manual. Trends Cell Biol.

Vrabioiu, A. M., and Mitchison, T. J. (2007). Symmetry of septin hourglass and ring structures. J. Mol. Biol. *372*, 37–49.

Wach, A. (1996). PCR-synthesis of marker cassettes with long flanking homology regions for gene disruptions in *S*. *cerevisiae*. Yeast *12*, 259–265.

Weirich, C. S., Erzberger, J. P., and Barral, Y. (2008). The septin family of GTPases: architecture and dynamics. Nat. Rev. Mol. Cell Biol. *9*, 478–489.

Wendland, J., Ayad-Durieux, Y., Knechtle, P., Rebischung, C., and Philippsen, P. (2000). PCR-based gene targeting in the filamentous fungus *Ashbya gossypii*. Gene *242*, 381–391.

Wendland, J., and Walther, A. (2005). *Ashbya gossypii*: a model for fungal developmental biology. Nat. Rev. Microbiol. *3*, 421–429.

Westfall, P. J., and Momany, M. (2002). Aspergillus nidulans septin AspB plays pre- and postmitotic roles in septum, branch, and conidiophore development. Mol. Biol. Cell *13*, 110–118.

Wightman, R., Bates, S., Amornrrattanapan, P., and Sudbery, P. (2004). In *Candida albicans*, the Nim1 kinases Gin4 and Hsl1 negatively regulate pseudohypha formation and Gin4 also controls septin organization. J. Cell Biol. *164*, 581–591.

Xie, Y., Vessey, J. P., Konecna, A., Dahm, R., Macchi, P., and Kiebler, M. A. (2007). The GTP-binding protein Septin 7 is critical for dendrite branching and dendritic-spine morphology. Curr. Biol. *17*, 1746–1751.