Spitzenkörper, Exocyst, and Polarisome Components in *Candida albicans* Hyphae Show Different Patterns of Localization and Have Distinct Dynamic Properties*†‡

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During the extreme polarized growth of fungal hyphae, secretory vesicles are thought to accumulate in a subapical region called the Spitzenkörper. The human fungal pathogen *Candida albicans* can grow in a budding yeast or hyphal form. When it grows as hyphae, Mlc1 accumulates in a subapical spot suggestive of a Spitzenkörper-like structure, while the polisome components Spa2 and Bud6 localize to a surface crescent. Here we show that the vesicle-associated protein Sec4 also localizes to a spot, confirming that secretory vesicles accumulate in the putative *C. albicans* Spitzenkörper. In contrast, exocyst components localize to a surface crescent. Using a combination of fluorescence recovery after photobleaching (FRAP) and fluorescence loss in photobleaching (FLIP) experiments and cytochalasin A to disrupt actin cables, we showed that Spitzenkörper-localized proteins are highly dynamic. In contrast, exocyst and polisome components are stably located at the cell surface. It is thought that in *Saccharomyces cerevisiae* exocyst components are transported to the cell surface on secretory vesicles along actin cables. If each vesicle carried its own complement of exocyst components, then it would be expected that exocyst components would be as dynamic as Sec4 and would have the same pattern of localization. This is not what we observe in *C. albicans*. We propose a model in which a stream of vesicles arrives at the tip and accumulates in the Spitzenkörper before onward delivery to the plasma membrane mediated by exocyst and polisome components that are more stable residents of the cell surface.

Polarized growth of fungi requires that a supply of secretory vesicles is delivered along cytoskeletal tracks to the site of cell expansion (for reviews, see references 13, 29, 30, and 31). Fusion of these membrane-bound vesicles with the plasma membrane allows the necessary expansion of the plasma membrane and releases the enzymes and raw materials for the synthesis of new cell wall material and the remodeling necessary to allow this newly synthesized material to be inserted into the existing cell wall. The process of polarized growth has been extensively studied in the budding yeast *Saccharomyces cerevisiae* and provides a model for studying the process in other fungi (for a review, see reference 20). Post-Golgi vesicles travel to sites of polarized growth along actin cables (23). Actin cables are nucleated at sites of polarized growth by the formin Bni1 facilitated by a multiprotein complex called the polisome, which consists of Spa2, Bud6, and Pea1 (5, 22, 24, 27). The motive force for vesicle transport is provided by Myo2, a class V myosin, complexed to its regulatory light chain Mlc1 (22, 26). At the plasma membrane, secretory vesicles dock with a second multiprotein complex called the exocyst before fusion with the plasma membrane (14, 15, 32, 33), mediated by v-SNARES on the vesicle and t-SNARES on the membrane.

The exocyst is an octomeric complex composed of Sec3, Sec5, Sec6, Sec8, Sec10, Sec15, Exo70, and Exo84 (21). It is thought that Sec3 and a portion of the Exo70 pool are localized at sites of polarized growth independently of the actin cytoskeleton (3, 6). The other exocyst subunits and the remainder of the Exo70 pool are thought to be transported to sites of polarized growth on secretory vesicles, where together with Sec3 and Exo70 they form the exocyst complex (3). Secretory vesicles exit the Golgi apparatus, travel toward sites of polarized growth, and dock with the exocyst by use of the Rab-type GTPase Sec4 in its GTP-bound form, which is activated by its GEF, Sec2 (12, 19, 35, 36). In the *S. cerevisiae* cell cycle, polarized growth is initially directed toward the bud tip in young buds (17). Growth subsequently becomes isotropic in larger buds before being directed toward the mother bud neck during cytokinesis at the end of the cell cycle. Accordingly, polisome and exocyst components localize to the tips of young buds (7, 27, 28).

The rate of hyphal tip extension is much greater than that of the growth of a yeast or pseudohyphal bud. In rich yeast extract-peptone-dextrose (YPD) medium, *Candida albicans* hyphae extend at the rate of 0.25 μm min⁻¹, compared to 0.0625 μm min⁻¹ in yeast buds and 0.125 μm min⁻¹ in pseudohyphal cells (P. Sudbery unpublished observations). In hyphae of filamentous fungi, a structure called a Spitzenkörper is present at the tip, which is rich in secretory vesicles (8, 9, 11, 29, 34). It is believed that the Spitzenkörper acts as a vesicle supply center (VSC) (1). This model proposes that the Spitzenkörper is maintained at a fixed distance from the hyphal tip. Vesicles radiate out in equal directions to fuse with the plasma membrane, so that more vesicles per unit area fuse with the hyphal tip itself than with other parts of the hyphae. Mathematical
modeling shows that this explains the distinctive shape of hyphal tips.

In order to investigate the mechanism of polarized growth in the hyphae of *Candida albicans*, we previously determined the localization of Mlc1-yellow fluorescent protein (YFP) and the polarisome components Bud6-YFP and Spa2-YFP (4). We found that in hyphae, polarisome components localized to a surface crescent, as they did in young yeast buds and the tips of elongated pseudohyphal buds. However, in hyphae Mlc1-YFP localized to a bright spot, which at least in some hyphae was clearly inside the tip, rather than at the surface, and which appeared spherical in three-dimensional reconstructions. We concluded that this represented a Spitzenkörper. In some hyphae Mlc1-YFP also localized to a surface crescent, similar to the pattern displayed by polarisome components. This observation suggested that the Spitzenkörper and polarisome were separate structures, both of which were present at hyphal tips, but that only the polarisome was present at the bud tips of pseudohyphae and yeast. Moreover, the dual localization of Mlc1-YFP to a crescent and a spot suggested that Mlc1 may be present in both structures.

While *S. cerevisiae* has proved to be an excellent model to investigate the molecular genetics of polarized growth, it is less optimal to study the spatial organization of the molecular components because polarized growth of the bud is restricted to a short period after bud emergence when the nascent bud is small. Thus, there has been little effort to investigate the fine details of the spatial organization of the different components of the polarized machinery beyond noting that they localize to sites of polarized growth. In this study we exploited the opportunities afforded by the continuous polarized growth of *C. albicans* hyphae to clarify the relationship between the Spitzenkörper, polarisome, and exocyst, which cooperate to mediate the extreme polarized growth of hyphae. We show that the vesicle-associated marker Sec4 also localizes to a Spitzenkörper-like structure, confirming the existence of a vesicle-rich area corresponding to a Spitzenkörper at the hyphal tip. We show that exocyst components such as Sec3, Sec6, Sec8, Exo70, and Exo84 localize to a surface crescent, so the exocyst, like the polarisome, is also a spatially separate structure from the Spitzenkörper. We used three independent strategies to investigate the dynamic properties of these structures. Fluorescence recovery after photobleaching (FRAP) was used to measure the rate at which new proteins arrived at the tip. Fluorescence loss in photobleaching (FLIP) was used to measure the rate at which proteins exited the tip. Cytochalasin A was used to disrupt actin cables, allowing the persistence of proteins at the tip to be measured after the supply of new proteins was blocked. In each case we found that Spitzenkörper components Sec4, Sec2, and Mlc1 were highly dynamic, while the polarisome component Spa2 was stable. Intriguingly, exocyst components showed intermediate dynamic properties, suggesting that they are delivered to the tip on vesicles but that not all vesicles carry a complement of exocyst components. We suggest that these data are consistent with a model in which a stream of vesicles arrives at the tip and accumulates in the Spitzenkörper before onward delivery to the plasma membrane mediated by exocyst and polarisome components that are more stable residents of the cell surface.

### MATERIALS AND METHODS

#### Strains

All strains were derived from BWPi7 (ura3ΔΔΔΔ hisG4/ΔΔΔΔ his1 his1 hisG4/ΔΔΔΔ his1). Genes encoding homologues of *S. cerevisiae* exocyst components were identified by two-way BLAST searches, the results of which were supported by annotation in the *Candida* genome database (www.candidagenome.org/). C-terminal fusions were generated as previously described using pFA-YFP plasmids carrying the appropriate URAs3, ARGS4, or HIS3 genes (10, 25). Full genotypes of the strains are provided in Table 1, and the oligonucleotides used are listed in Table S1 in the supplemental material.

#### Growth conditions

Hyphal growth was induced by growing yeast cells at 25°C overnight to saturation in YEPD (2% glucose, 2% Difco Bacto peptone, and 1% Difco Bacto yeast extract plus 80 mg ml⁻¹ uridine). The stationary-phase culture was washed with distilled water and inoculated at a 1:20 dilution into synthetic defined (SD) medium (consisting of 0.67% yeast nitrogen base [Difco], 2% [wt/vol] glucose, and 80 mg ml⁻¹ each of histidine, uridine, and arginine in phosphate-buffered saline [PBS], pH 7.0) plus 20% calf serum (Sigma-Aldrich) and incubated at 37°C. Under these conditions, hyphae evaginate after 35 to 40 min. For live-cell imaging in FRAP and FLIP experiments, cells were grown on agar pads incorporating SD medium as described previously (2).

Cytochalasin A (Sigma-Aldrich) was used at working concentration of 10 μM from a stock concentration of 1 mg ml⁻¹ in dimethyl sulfoxide (DMSO). Typically, 5 μl of the stock solution was added to 1 ml of cells. Control experiments showed that the DMSO alone had no effect on hyphal growth or localization of the fluorescent proteins.

#### Microscopy

Images of exocyst components fused to YFP and green fluorescent protein (GFP)-Sec4 localization were made using a Delta Vision Spectris 4.0 RT microscope (Applied Precision Instruments, Seattle, WA) using an Olympus 100x UPlanApo NA 1.35 lens (Olympus, Tokyo, Japan). Images were acquired and deconvolved with Softworx software. To visualize the cell outline in fluorescent images, cells were counterstained with 1 μg/ml calcofluor white (Fluorescent Brightener 28; Sigma-Aldrich, St. Louis, MO).

All FRAP and FLIP experiments were carried out on hyphae actively growing on the surface of an agar pad in a concavity slide maintained at 37°C by a heated concave heating device. The hyphal tips were illuminated with blue light to bleach specific regions. To quantify FRAP and FLIP images, the image acquisition parameters were calibrated using this system.
TABLE 2. *C. albicans* homologues of *S. cerevisiae* exocyst components

<table>
<thead>
<tr>
<th><em>S. cerevisiae</em> protein, ORF</th>
<th>Best hit in <em>C. albicans</em> genome</th>
<th>E value&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Assigned gene name in CGD&lt;sup&gt;b&lt;/sup&gt;</th>
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</thead>
<tbody>
<tr>
<td>Sec3, YER008C</td>
<td>Orf19.2911</td>
<td>2.5e–18</td>
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<tr>
<td>Sec6, YIL066C</td>
<td>Orf19.5463</td>
<td>1.5e–47</td>
<td>SEC6</td>
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<tr>
<td>Sec8, YPR055W</td>
<td>Orf19.3647</td>
<td>1.2e–93</td>
<td>SEC8</td>
</tr>
<tr>
<td>Exo70, YIL085W</td>
<td>Orf19.6512</td>
<td>3.5e–48</td>
<td>EXO70</td>
</tr>
<tr>
<td>Exo84, YBR102C</td>
<td>Orf19.135</td>
<td>3.2e–30</td>
<td>EXO84</td>
</tr>
</tbody>
</table>

<sup>a</sup> E values are from BLAST searches using the *S. cerevisiae* protein sequence to probe the *C. albicans* genome.

<sup>b</sup> CGD, *C. albicans* genome database (www.candidagenome.org).

stage and environment chamber (Pecon GmbH, Erbach, Germany). FRAP and FLIP experiments were carried out using a Zeiss LSM 510 Meta confocal microscope running release 4.0 SP2 software. The 514-nm HFT laser and the long-pass 505 filter were used to image GFP. Photobleaching was carried out using the argon/2 laser, which was set up with tube power at 6.0 Å and used at 45% of the total power and with the region of interest (ROI) positioned to expose the hyphal tip of the bleached cell (FRAP) or a strip 1 μm wide located approximately 3 μm behind the tip (FLIP). A nonbleached tip in the same field served as a control for bleaching due to image acquisition in the recovery period. Average image intensity was measured using LSM software for the experiments shown in Fig. S3 in the supplemental material and using NIH Image J for the experiments shown in Fig. 3 and 4 and in Fig. S2 in the supplemental material. Raw values were subject to a subtraction of background. Values were then corrected for bleaching due to image acquisition in the recovery period according to the proportional loss of intensity in the control tip at the same time point, which was set to 100% prebleach. For each time point the fraction of fluorescence recovered (FRAP) or lost (FLIP) was calculated according to the formula \( (I_t - I_0)/(I_0 - I_t) \times 100\% \), where \( I_t \) = time in seconds, \( I_0 \) = intensity prebleaching, \( I_t \) = intensity immediately postbleach, and \( I_t \) = intensity at time \( t \). Prism software (GraphPad Software Inc.) was then used to fit the best curve to a plot of time versus fraction fluorescence recovered. In all experiments analyzed, the best fit was to a nonlinear curve with a one-phase exponential. The same software was used to generate the bar chart and carry out \( t \) tests for pairwise comparisons shown in Fig. 3.

RESULTS

Exocyst components localize to a surface crescent, but Sec4 localizes to a Spitzenkörper-like structure. We carried out BLAST searches to identify *C. albicans* orthologues of *S. cerevisiae* Sec3, Sec6, Sec8, Exo70, and Exo84 proteins (Table 2). Except for Sec3, all proteins identified were of similar length, and the similarity extended across the full length of the protein. Although the similarity was less clear in the case of Orf19.2911, the putative *C. albicans* Sec3 orthologue, it has been shown that this gene can functionally complement an *S. cerevisiae sec3* mutant and localizes to hyphal tips (18). We generated C-terminal YFP fusions of each of these *C. albicans* exocyst orthologues. In each case the exocyst component clearly localized to a surface crescent in the tips of essentially all hyphae examined. Figure 1 shows the pattern of localization of representative hyphal tips for each exocyst protein. Figure S1 in the supplemental material shows a field of cells expressing Sec3-YFP, with each tip enlarged to show the consistency of the localization pattern. A similar consistency was seen in the localization patterns of the other exocyst components. Although there were bright spots of fluorescence within the crescent, these were clearly qualitatively different from the spot-like localization of Sec4, which we discuss further below. We also obtained similar results for a putative Sec15 homologue (R. Lane and P. Sudbery, unpublished observations). Thus, like the polarisome (4), exocyst components localize to a surface crescent and not the Spitzenkörper.

An N-terminal GFP-Sec4 fusion has previously been shown to localize to hyphal tips, but no attempt was made to distin-
guish whether its localization was to a spot or crescent (18). We have reexamined this strain (kindly provided by Yue Wang) and found that GFP-Sec4 localizes to a spot, rather than a crescent, in essentially all hyphae examined. (Fig. 2A; see Fig. S1 in the supplemental material). Since Sec4 is a recognized secretory vesicle-associated marker (19), this supports the notion that the spherical structure previously observed in *C. albicans* hyphae is indeed a vesicle-rich Spitzenkörper. We have also obtained a similar result with Sec2, the GEF for Sec4 (2). As pointed out above, bright spots were sometimes visible in the crescent-like localization of exocyst components. However, the localization of GFP-Sec4 was qualitatively different. Figure 2B shows a montage of the individual Z-planes of GFP-Sec4 and Exo84-YFP. The large GFP-Sec4 spot is clearly visible.
FIG. 3. Fluorescence recovery of Spitzenkörper, exocyst, and polarisome components. Strains expressing the indicated proteins fused to YFP or GFP (Sec4) were grown on surface agar concavity slides under hypha-inducing conditions for 90 min. A minimum of 10 hyphal tips of each strain were bleached with a Zeiss LSM 510 as described in Materials and Methods, and images were recorded immediately before bleaching (prebleach), 1.5 s after bleaching (postbleach), and at 30-s intervals after bleaching (+30s). (A) The average fluorescence intensity was measured in each tip, and the percentage of fluorescence lost by bleaching that was recovered after 30 s was calculated according to the formula $(I_{30} - I_{b})/(I_0 - I_{b}) \times 100$, where $I_{30}$ = intensity at 30 s after bleaching, $I_b$ = postbleach intensity, and $I_0$ = prebleach intensity. Error bars are standard errors of the means. Results of pairwise $t$ tests are shown above the brackets. *** $P = 0.0001$; ** $P = 0.0014$; ns, difference not significant ($P = 0.1$). (B) Representative images of individual hyphal tips at prebleach, postbleach, and +30 s. Scale bar, 2 μm.
throughout the Z-stack, whereas the bright spots in Exo84-YFP are confined to individual focal planes and the surface crescent.

Similar investigations with Ashbya gossypii found that the pattern of localization of exocyst proteins and the polarisome protein Spa2 depended upon hyphal extension rates (16); in fast-growing hyphae these proteins localized to a spot, while in slow-growing hyphae they localized to a surface crescent. The observation that the pattern of localization of exocyst and polarisome components is dependent on growth rate in A. gossypii raises the question of whether the different patterns of localization that we have observed in C. albicans may simply be the result of variation in growth rates between strains carrying different fluorescent protein fusions. This question may be addressed by determining the pattern of localization of Spitzenkörper, polarisome, or exocyst components in the same cell, thus removing growth rate as a variable. Indeed, we previously showed that Spa2-YFP localized to a crescent and Mlc1-cyan fluorescent protein (CFP) to a Spitzenkörper in the same hypha (4). However, the quantum efficiency of CFP when fused to C. albicans proteins is very low, making visualization of the target protein difficult. The spectral characteristics of YFP and GFP prevent the signals being distinguished by the Delta Vision wide-field epifluorescence microscope that we routinely use. However, YFP and GFP signals can be distinguished with a Zeiss laser scanning META confocal microscope using appropriate lasers for excitation and setting appropriate limits to the adjustable emission channels (see the legend Fig. 2). By this means we visualized Exo70-YFP and GFP-Sec4 in the same cell and confirmed that Exo70-YFP localized to a crescent and GFP-Sec4 to a spot (Fig. 2C).

FRAP experiments show that Spitzenkörper, exocyst, and polarisome components have different dynamic properties. We carried out fluorescence recovery after photobleaching (FRAP) experiments to investigate whether the different patterns of localization of exocyst, polarisome, and Spitzenkörper components are reflected in different dynamic properties. In such experiments the rate of fluorescence recovery, measured by the time constant \( \tau \), reports the rate at which new fluorescent components replace bleached components (the smaller the value, the more rapid the recovery). When the recovery curve reaches a plateau, the fraction of the original fluorescence regained is a measure of the fraction that is mobile. The use of FRAP was complicated by the time span of several minutes that it took for exocyst components to recover, while Spitzenkörper components recovered within a single minute. To accommodate these different time scales, we acquired post-bleach images every 5 s for GFP-Sec4, Mlc1-YFP, and Spa2-YFP (see Fig. S3 in the supplemental material) and every 30 s for GFP-Sec4 Mlc1-YFP, Sec8-YFP, Sec2-YFP, Bud6-YFP, Sec3-YFP, and Spa2-YFP (see Fig. S3 in the supplemental material). The derived values for \( \tau \), half-time \( (t_{1/2}) \), and mobile fraction (see Table S2 in the supplemental material) indicate that Sec4 and Mlc1 are most dynamic, the exocyst components were less dynamic, and the polarisome component Bud6 was least dynamic.

The time span of several minutes for exocyst components to show full recovery made it difficult to generate enough reliable replicates for statistical analysis, because the hyphal tip would frequently grow in the Z dimension, rendering it out of focus in the postbleach images. Refocusing during this period would distort the results, as the additional image acquisition required would itself cause bleaching of the fluorophore being measured. (In C. albicans, YFP and GFP are very sensitive to photobleaching.) Although it was possible to calculate \( \tau \) values from these data, the confidence limits were large, which made comparisons between strains problematic (see Table S2 in the supplemental material). Since we had established that exocyst components did eventually recover their full fluorescence (see Table S2 in the supplemental material), the fraction recovered after 30 s is dependent only on the recovery rate and not on the mobile fraction. Thus, we used Sec4 as a benchmark to compare the fraction of fluorescence recovered in a single image acquired after 30 s in a minimum of 10 replicate hyphal tips (20 in the case of Mlc1-YFP). Moreover, the single postbleach image acquired in this protocol has the additional benefit that a bleaching correction was not necessary. The results are shown in Fig. 3A. Consistent with the previous experiments, GFP-Sec4 showed the most rapid recovery. The next-highest rate was shown by Mlc1-YFP and Sec2-YFP, both of which, like Sec4, localize to the Spitzenkörper. Although the difference in the rates of recovery for Sec2 and Sec4 fell short of statistical significance in a two-tailed \( t \) test (\( P = 0.1 \)), the pattern was reproduced in previous experiments and is likely to be real. The exocyst components, Exo70-YFP, Sec3-YFP, Exo84-YFP, Sec8-YFP, and Sec6-YFP, formed a group with intermediate recovery rates. The difference in the rates of recovery of Mlc1-YFP and Exo70-YFP, the most dynamic exocyst component, was significant (\( P = 0.0014 \)). The polarisome component Spa2-YFP showed the lowest recovery rate; the difference between the rates of recovery of Spa2-YFP and Sec8-YFP was highly significant (\( P = 0.0001 \)). Figure 3B shows images of hyphal tips prebleach, postbleach, and after 30 s of recovery. GFP-Sec4, Mlc1-YFP, Exo70-YFP, and Spa2 are shown to represent each class of protein described above with respect to the rate of recovery. Taken together, these results show that the different patterns of localization of vesicle-associated proteins that localize to the Spitzenkörper, such as Sec4, Mlc1, and Sec2, and exocyst and polarisome components is reflected in the different dynamic properties of these proteins.

The Spitzenkörper is in dynamic equilibrium between vesicles entering and leaving. FRAP experiments determine the rate at which a bleached protein is exchanged for an unbleached protein. Thus, in the present context, these experiments measure the rate at which new, unbleached proteins arrive at the tip. The rate at which proteins are lost from the tip can be measured by a fluorescence loss in photobleaching (FLIP) experiment, in which a strip just behind the tip is continually bleached, ablating the fluorescence of proteins before they arrive at the tip. Fluorescence declines because fluorescence on vesicles leaving the tip is not replaced by the fluorescence of newly arriving proteins. GFP-Sec4 or Spa2-YFP was grown under hypha-inducing conditions and repeatedly bleached at 1-s intervals in a strip just behind the tip. To correct for bleaching during image acquisition after each bleaching event, a field was chosen with two hyphal tips, one of which served as a control. The results are expressed as the fraction of initial fluorescence remaining in the bleached hyphae normalized to the fraction of initial fluorescence remaining in the control tip (Fig. 4). In the case of GFP-Sec4, fluo-
Fluorescence rapidly declined. The time constant (τ) of the exponential decline was 11.7 s (95% confidence interval, 8.2 to 20.3 s). This value is similar to the τ value of 10.6 s (95% confidence interval, 7.5 to 18.0 s) calculated in FRAP experiments (see Table S2 and Fig. S2 in the supplemental material), suggesting that Sec4 in the Spitzenkörper is in a state of dynamic flux, with a constant stream of incoming molecules replacing molecules that exit. In contrast, the fluorescence of Spa2-YFP showed only a modest decline during the course of the FLIP experiments, suggesting that Spa2 is a stable component of the polarisome at the cell surface.

Spitzenkörper components, but not exocyst or polarisome components, depolarize rapidly upon disruption of actin cables. The FRAP and FLIP experiments described above suggest that Sec4 is more dynamic than the exocyst components. If GFP-Sec4 and exocyst components are delivered to the tip along actin cables, then Sec4 should depolarize faster than exocyst components upon disruption of actin cables. To test this prediction, we used cytochalasin A, which disrupts actin cables and leads to tip swelling after 15 min. After the addition of cytochalasin A to liquid cultures, we monitored the persistence of polarized Spitzenkörper, polarisome, and exocyst components. There was a clear difference in the rates of depolarization of Spitzenkörper components and of exocyst and polarisome components. GFP-Sec4 depolarized within 90 s upon addition of cytochalasin A, and the fluorescence dispersed into bright punctate spots distributed throughout the germ tube mother cell (Fig. 5A). The value of 90 s is a maximum figure, representing the shortest time taken after addition of the drug to a liquid culture to withdraw a sample, prepare and mount a slide, and focus and record an image. Mlc1-YFP (Fig. 5B and C) and Sec2-YFP (data not shown) also depolarized rapidly, although after 3 min some polarized material remained, consistent with the slightly slower kinetics of FRAP. In the case of Mlc1-YFP, a faint surface crescent remained in some hyphae (Fig. 5C), which persisted for up to 15 min; this
may represent Mlc1-YFP present in the polarisome as we previously reported.

In contrast, exocyst and polarisome components were all still fully polarized after 5 min. Sec3-YFP is shown as an example in Fig. 6A. In the case of Sec3, the fluorescence eventually dispersed into punctate spots by 15 min (Fig. 6B). In the case of Spa2, the fluorescence was still clearly visible after 15 min as the tip started to swell (Fig. 6D). In the case of other exocyst components, the fluorescence slowly faded at the tip, but the crescent was still visible as the tip started to swell after 15 min (e.g., Sec8-YFP) (Fig. 6C). Thus, consistent with the FRAP and FLIP experiments, the Spitzenkörper components are in a state of rapid dynamic flux, while the polarisome and Spitzenkörper components are more stable.

**DISCUSSION**

We previously showed that at the tip of *C. albicans* hyphae, Mlc1-YFP localized to a spot while the polarisome components Spa2 and Bud6 localized to a crescent. We suggested that Mlc1 was localizing to a Spitzenkörper and that this was a spatially distinct structure from the polarisome. Here we extend this work by showing that GFP-Sec4 also localizes to a spot. We describe elsewhere a similar localization pattern for Sec2, the GEF for Sec4 (2). Not only do these observations confirm the existence of a Spitzenkörper in *C. albicans* hyphal tips, but they also show that the Spitzenkörper is indeed rich in secretory vesicles, as suggested by electron microscope images of the Spitzenkörper in other fungal hyphae. We further show that exocyst components localize to a surface crescent, suggesting that the exocyst is a structure that is spatially distinct from the Spitzenkörper. It should be noted that in this work we did not formally assess the functionality of the exocyst-YFP fusions. However, in both *S. cerevisiae* and *A. gossypii*, all of these fusions have been shown to be functional (16). Moreover, the similar localization patterns of all the exocyst components provide additional confidence that the observations are reliable.

Localization of exocyst and polarisome components has been reported for *A. gossypii* (16). Upon spore germination, hyphae initially grow at a lower rate than is attained in longer
It was found that the localization pattern of exocyst components depended on the growth rate. In the slow growth phase, these components localize to a surface crescent as we find in *C. albicans* hyphae. However, at a higher growth rate, both the exocyst and Spa2 were found to localize to the Spitzenkörper. These observations can be reconciled with our observations by considering that hyphal growth in *C. albicans* is analogous to the slow mode of hyphal growth in *A. gossypii*, so that in both cases exocyst components and Spa2 localize to a crescent. According this scenario, Mlc1, Sec4, and Sec2 would localize to the Spitzenkörper in slow-growing *A. gossypii* hyphae. Unfortunately, the localization of these components was not reported to allow this prediction to be tested.

Having established that Sec4, Mlc1, and Sec2 localize to a spot whereas polarisome and exocyst components localize to a crescent, we next investigated whether these differences in spatial organization were reflected in different dynamic properties. Three independent experimental strategies confirmed that this was indeed the case. The reliability of conventional FRAP recovery curves was compromised by the time span of several minutes required for full recovery of exocyst and polarisome components and the broad confidence limits on the
estimations of $\tau$. Although we generated such curves and report them in the supplemental material, we consider the fraction of fluorescence recovered by 30 s to be a more reliable guide. It is important to note that the full FRAP curves show that exocyst components did eventually recover 100% of the original fluorescence so that the fraction recovered by 30 s is not limited by the size of the mobile fraction. Both types of FRAP experiment led to the same conclusion that the Spitzenkörper components GFP-Sec4, Mlc1, and Sec2 were more dynamic than the exocyst components, which in turn were more dynamic than the polarisome components Spa2 and Bud6. Among the Spitzenkörper components, GFP-Sec4 appeared to be more dynamic than Mlc1 or Sec2. An explanation for this difference may be that Sec2 may detach from the vesicles after activating Sec4. Similarly, Mlc1 may be released from the Myo2-secretory vesicle complex after it arrives at the Spitzenkörper. In contrast Sec4 may remain on the vesicle until its fusion with the plasma membrane.

The full FRAP curves (see Fig. S2 and S3 and Table S2 in the supplemental material) generated a $\tau$ value for GFP-Sec4 (10.7 s) which is very similar to the value reported by Boyd et al. for Sec4 in S. cerevisiae. Interestingly, in that study Sec4 also showed a higher recovery rate than exocyst components (3). However, that study reported $\tau$ values in the range of 15 to 20 s for exocyst components other than Sec3 and Exo70, which is considerably faster than we observed. Interestingly, the mobile fraction for these exocyst components in S. cerevisiae was reported to be on the order of 40%, while we observed that the mobile fraction was close to 100%. An explanation for these differences is that in C. albicans exocyst components can be recruited from distant hyphal locations, so that there is a greater reservoir of unbleached components, but it takes longer for them to be recruited. Sec3 has been reported to be recruited independently of the actin cytoskeleton (6), and Boyd et al. (3) reported a slow recovery time for this exocyst subunit. Moreover, Boyd et al. reported a two-phase recovery for Exo70 and suggested that a fraction of the Exo70 pool was recruited independently of actin cables. We did not observe that Sec3 was less dynamic than the other exocyst components. We also observed a one-phase recovery of Exo70-YFP. Thus, in C. albicans Exo70 and Sec3 appear to be delivered along actin cables. A similar conclusion was reported for these proteins in A. gossypii (16).

In addition to the FRAP experiments, we investigated the dynamic properties in two further ways. First, we used FLIP experiments, which determine the rate at which fluorescence declines when the fluorescence of incoming GFP-Sec4 proteins is ablated by bleaching before they arrive at the tip. We interpret this as a measure of the rate at which vesicles exit the tip. However, an alternative explanation is that GFP-Sec4 detaches from the vesicle once it arrives at the Spitzenkörper. Regardless of which interpretation is correct, the FLIP experiment provides an independent measure of the dynamic properties of Sec4. Importantly, the $\tau$ value of the recovery in FRAP experiments (10.6 s; 95% confidence limits, 7.5 s to 18.0 s) was very similar to the $\tau$ value of the exponential in the FLIP experiments (11.7 s; 95% confidence limits, 8.2 s to 20.3 s). Taken together, the FRAP and FLIP experiments show that the flux of Sec4 molecules entering and leaving the Spitzenkörper is highly dynamic. Spa2 was much less dynamic in the FLIP experiments, confirming the results of the FRAP experiments.

Finally, we interrupted the supply of molecules to each tip by disrupting actin cables. Again we observed that Sec4, Sec2, and Mlc1 rapidly depolarized upon cytochalasin A treatment. Sec2 was completely depolarized within 90 s. Sec2 and Mlc1 were largely depolarized within 3 min, although some polarized material remained. Interestingly, the remaining polarized Mlc1 protein was distributed in a surface crescent which may represent Mlc1 in the polarisome, as we previously reported (4). In contrast, the polarisome and exocyst components were much more stable. Spa2 remained polarized even as the tip started to swell after 15 min. Exocyst components were also still polarized at this time, although the intensity of the fluorescence had diminished. A model for hyphal growth. The current paradigm for polarized growth in S. cerevisiae considers that exocyst subunits travel to the cell surface on secretory vesicles. (3). If this was the case in C. albicans, we would expect that the pattern of localization and dynamic properties of vesicle-associated proteins such as Sec4 and Sec2 would be similar to that of exocyst components. The data presented here provide compelling evidence that this is not the case. Sec4, Sec2, and Mlc1 are clearly transported to the tip on actin cables, where they accumulate in the Spitzenkörper. They are highly dynamic, with a flux of incoming molecules balanced by an outflow, presumably to the cell surface. Exocyst and polarisome components are stably located in a surface crescent. They may be transported to the cell surface on actin cables, but our data leave this question open, as it is possible that the slow depolarization upon cytochalasin A treatment is a secondary consequence of the disruption of actin cables. If they are transported on actin cables, the difference in their dynamic properties compared to those of Spitzenkörper components suggests that only a frac-
tion of vesicles carry exocyst components and that once delivered to the cell surface they remain to act as docking platforms for other incoming vesicles.

Our results suggest that a modified form of the vesicle supply model operates in *C. albicans* hyphae; this model which attempts to explain why proteins such as Sec4 are localized to an apical spot whereas exocyst and polarisome components localize to a surface crescent (Fig. 7). The essential feature of the model is that secretory vesicles marked by Sec4, Sec2, and Mlc1 arrive in a constant stream at the Spitzenkörper along actin cables. The Spitzenkörper is in dynamic balance, with vesicles leaving at the same rate at which they arrive. Post-Spitzenkörper vesicles are delivered to the cell surface along actin cables nucleated by the polarisome located at the cell surface. At the cell surface, they are tethered by exocyst structures which may be carried on the vesicle or may have been left behind after delivery by other vesicles.

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