In *Candida albicans*, phosphorylation of Exo84 by Cdk1-Hgc1 is necessary for efficient hyphal extension

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**ABSTRACT** The exocyst, a conserved multiprotein complex, tethers secretory vesicles before fusion with the plasma membrane; thus it is essential for cell surface expansion. In both *Saccharomyces cerevisiae* and mammalian cells, cell surface expansion is halted during mitosis. In *S. cerevisiae*, phosphorylation of the exocyst component Exo84 by Cdk1-Clb2 during mitosis causes the exocyst to disassemble. Here we show that the hyphae of the human fungal pathogen *Candida albicans* continue to extend throughout the whole of mitosis. We show that CaExo84 is phosphorylated by Cdk1, which is necessary for efficient hyphal extension. This action of Cdk1 depends on the hyphal-specific cyclin Hgc1, the homologue of G1 cyclins in budding yeast. Phosphorylation of CaExo84 does not alter its localization but does alter its affinity for phosphatidylinerine, allowing it to recycle at the plasma membrane. The different action of Cdk1 on CaExo84 and ScExo84 is consistent with the different locations of the Cdk1 target sites in the two proteins. Thus this conserved component of polarized growth has evolved so that its phosphoregulation mediates the dramatically different patterns of growth shown by these two organisms.

**INTRODUCTION**
*Candida albicans* can grow in budding yeast both pseudohyphal and true hyphal forms (Sudbery et al., 2004). This morphological plasticity is essential for virulence of this major human fungal pathogen. In the hyphal state, growth is highly polarized to the tip (Soll et al., 1985). Such polarized growth requires a constant supply of secretory vesicles that fuse with the plasma membrane at the tip (Sudbery, 2011; Caballero-Lima et al., 2013). These vesicles are necessary for cell surface expansion, as they supply the additional membrane necessary for expansion of the plasma membrane. Vesicles also deliver the means to synthesize new cell wall, such as integral membrane glucan synthases, which manufacture β1,3-glucan and β1,6-glucan, chitin synthases, cell wall–remodeling enzymes, and the mannosylated proteins that form the outer layer of the cell wall (Klis et al., 2006). Studies in the model yeast *Saccharomyces cerevisiae* showed that before fusion, secretory vesicles must be tethered to the plasma membrane by an evolutionarily conserved, multiprotein complex called the exocyst. This complex consists of eight subunits: Sec3, Sec5, Sec6, Sec8, Sec10, Sec15, Exo70, and Exo84 (Terbush and Novick, 1995; Terbush et al., 1996; Brennanwald and Rossi, 2007; He and Guo, 2009; Heider and Munson, 2012). In *S. cerevisiae* these proteins localize to sites of polarized growth such as small buds and the site of septum formation. Sec3, as well as part of the Exo70 pool, is believed to localize independently of the cytoskeleton (Finger et al., 1998; Boyd et al., 2004). Exo70 is an effector of and interacts with the Rho-type GTPases Rho3 and Cdc42, and Sec3 interacts with Rho1 (Adamo et al., 1999; Guo et al., 2001; Zhang et al., 2001; Roumanie et al., 2005; He et al., 2007b; Wu et al., 2010). Both Exo70 and Sec3 interact with phosphoinositol 4,5-phosphate (He et al., 2007a; Zhang et al., 2008). The remaining exocyst components are believed to arrive on incoming vesicles (Boyd et al., 2004).
In C. albicans, exocyst subunits localize to a surface crescent at the hyphal tip (Jones and Sudbery, 2010). In contrast, vesicle-associated proteins such as Sec4, Sec2, and Mlc1 localize to a subapical spot that is clearly distinct from the surface crescent of the exocyst and is reminiscent of the Spitzenkörper long known to drive the polarized growth of other filamentous fungi (Crampin et al., 2005; Bishop et al., 2010; Jones and Sudbery, 2010). These different localization patterns suggest that at least during the polarized growth of hyphae, the exocyst is already present on the plasma membrane to tether the vesicles as they arrive. Such a notion is supported by fluorescence recovery after photobleaching (FRAP) and fluorescence loss in photobleaching (FLIP) experiments, which show that Sec2, Sec4, and Mlc1 are much more dynamic than exocyst components. Furthermore, disruption of the actin cytoskeleton leads to immediate dispersal of Sec4 from the Spitzenkörper, whereas exocyst components disperse more slowly from the surface crescent (Jones and Sudbery, 2010).

As well as controlling progress through the cell cycle, it is becoming increasingly apparent that the cyclin-dependent kinase Cdk1 (Cdc28) controls cell growth in S. cerevisiae (McCusker et al., 2007; Goranov and Amon, 2010). In C. albicans the cyclin Hgc1 is specifically induced during hyphal growth and is necessary for normal hyphal morphology and growth (Zheng et al., 2004). Hgc1 is the homologue of the Cln1/2 G1 cyclin pair of S. cerevisiae. Cells lacking Hgc1 are unable to maintain hyphal growth. When induced to form hyphae, such cells evaginate a normal germ tube, but this quickly swells, especially after the formation of the first septin ring. A number of targets for Cdk1-Hgc1 have been elucidated, such as the Cdc42 GAP Rga2, Sec2, the septin Cdc11, and the transcription factor Elf1 (Sinha et al., 2007; Zheng et al., 2007; Wang et al., 2009; Bishop et al., 2010). G2 cyclins in C. albicans are represented by Cib2, the orthologue of S. cerevisiae Cib2, and Cib4, the orthologue of S. cerevisiae Cib4. Cib2 is essential; deletion of Cib2 using the MET3-regulated promoter results in cell cycle arrest with highly elongated buds, whereas ΔCib4 cells are viable but show constitutive pseudohyphal growth (Bensen et al., 2005).

In S. cerevisiae it was recently shown that phosphorylation of the exocyst subunit Exo84 by Cdk1-Cib2 causes cell growth to cease just before the metaphase-anaphase transition (Luo et al., 2013). Exo84 plays a key role in the assembly of the exocyst. Its phosphorylation by Cdk1-Cib2 causes the exocyst complex to disassemble, blocking exocytosis and cell surface expansion. ScExo84 and CaExo84 show similar domain organization consisting of a coiled-coiled Vps51 domain that in Vps51 is required for formation of the VFT docking complex mediating the fusion of late endosomes with the Golgi (Siniassoglou and Pelham, 2002), a central pleckstrin homology (PH) domain, and a C-terminal, rod-like domain composed of multiple α-helices that is responsible for the interaction with other members of the exocyst complex (Figure 1A). Luo et al. studied the effects of Cib2-Cdk1 phosphorylation of ScExo84 at two full Cdk1 consensus target sites (S/T/P.X/R/K) and three minimal target sites (S/T/P; Luo et al., 2013). In view of the observation that phosphorylation of ScExo84 disrupts interaction with other exocyst subunits, it is significant that one of the full consensus sites in ScExo84 is located within the C-terminal interaction domain (S716). Of interest, there is no corresponding site in the C-terminal domain of CaExo84 (Figure 1, A and B). However, CaExo84 contains a full site within the PH domain (S384), which is absent in ScExo84 (Figure 1, A and B). Two additional full sites are located in CaExo84 at S256 and T488, which a Clustal alignment suggests may correspond to S291 and T496, respectively, in ScExo84 (Figure 1B). Modeling of CaExo84 shows that these sites are located on either side of the PH domain in the folded protein (Figure 1A).

Although cessation of cell growth during mitosis is a common feature among eukaryotic cells, we show here that C. albicans hyphae continue to grow throughout mitosis and cytokinesis. Moreover, localization of both the Spitzenkörper and the exocyst is unaltered throughout mitosis. However, we show that CaExo84 is phosphorylated by CaCdk1 in a hyphal-specific manner and that this phosphorylation is dependent on the hyphal-specific Hgc1 cyclin rather than a G2 cyclin. Moreover, in direct contrast to budding yeast, this phosphorylation of Exo84 is essential for rapid polarized growth of hyphae and normal hyphal morphology. Our study suggests that a role of phosphorylation is to dissociate Exo84 from the plasma membrane by lowering its affinity for phosphatidylinerine.

**RESULTS**

*C. albicans* hyphae continue to grow throughout mitosis and cytokinetic ring contraction

To investigate whether *C. albicans* hyphae continue to extend throughout mitosis, we constructed strains in which the nucleolar protein Nop1 was fused to yellow fluorescent protein (YFP) to track the nucleus. In addition, we also fused either Mlc1 or Exo70 to YFP. Initially, we followed hyphal growth throughout two complete cell cycles in the strain expressing Nop1-YFP and Mlc1-YFP after hyphae were induced from stationary-phase, unbudded yeast cells. We recorded images at 60-s intervals (Supplemental Movie S1). As visualized by Nop1-YFP, the nucleus migrates out of the mother cell and divides within the germ tube, as reported previously (Sudbery, 2001). This is followed 20 min after anaphase by contraction of the cytokinetic ring visualized by M1c1-YFP. A second anaphase occurs after 30 min, which in turn is followed by cytokinetic ring contraction 20 min later. Throughout the whole of these two cycles the hyphae extend at a constant rate of 0.33 μm/min (Supplemental Figure S1A). Close examination around the time of the two mitoses and the two contractions of the cytokinetic ring showed that hypha growth continued to extend during these events at a rate consistent with the long-term average rate of hyphal extension (Supplemental Figure S1, B and C).

It may be argued that the cessation of growth occurs on a shorter time scale than the 60-s intervals between frames in Supplemental Movie S1. We therefore investigated the growth of cells over shorter time scales. Moreover, it was of interest to visualize the Spitzenkörper, as this is a sensitive indicator of polarized growth of fungal hyphae, and the exocyst to discover whether it is dispersed as it is in S. cerevisiae during metaphase. We therefore recorded images over 10-s intervals throughout mitosis for the Nop1-YFP Mlc1-YFP strain and 20-s intervals for the Nop1-Exo70 strain. These experiments were complicated by the extreme sensitivity of mitosis to the excitation light; for this reason the frame rate was increased to 20 s for the Nop1-YFP Exo70 YFP strain. The full movies are Supplemental Movies S2 and S3 for Mlc1-YFP Nop1-YFP and Nop1-YFP Exo70-YFP, respectively. Consecutive frames spanning a mitosis are shown in Figure 2 for Mlc1-YFP Nop1-YFP and in Supplemental Figure S2 for Nop1-YFP Exo70-YFP. Of interest, in the Nop1-YFP Mlc1-YFP strain a pair of satellite spots appeared on either side of the nucleus just before anaphase, which may represent recruitment of myosin to the spindle pole body during metaphase. Evidence for a myosin role in the mitotic spindle has recently been reviewed (Sandquist et al., 2011). Figure 2A shows that growth continued throughout mitosis, so that the tip extended by 0.8 μm in the 140-s of the movie shown in Figure 2A, consistent with the long-term average growth rate of
FIGURE 1: Domain organization and Cdk1 target sites of ScExo84 and CaExo84. (A) Domain organization and Cdk1 target sites. The VPS51 domain was identified in both CaExo84 and ScExo84 using the Pfam database (Punta et al., 2012). The PH domain and the C-terminal interaction domain were identified by threading as described in Materials and Methods. Studied sites refer to this article for CaExo84 and Luo et al. (2013) for ScExo84. The folding structure of the PH domain predicted by threading is shown in purple. Phosphorylation of potential Cdk1 target sites is shown by the yellow spheres. S284 lies within a protruding loop of the PH domain, and S256 and T488 are located at the boundaries of the PH domain. (B) Clustal alignments of sequences surrounding full Cdk1 sites in CaExo84. Color code follows standard Clustal X code: amino acids with similar properties are coded with the same color; the depth of the color denotes the degree of similarity, so that identical residues have the deepest color.
The Spitzenkörper remained in place before, during, and after mitosis, and the intensity of Mlc1-YFP did not diminish throughout metaphase and anaphase. This suggests that vesicle traffic into the Spitzenkörper and from the Spitzenkörper to the cell surface continued throughout mitosis. Supplemental Figure S2 shows that throughout mitosis the exocyst component Exo70 also remained as a crescent at the hyphal tip and that the hyphal tip continued to extend. Thus hyphal extension and polarized exocytosis continue throughout mitosis.

Exo84 is phosphorylated by Cdk1 in a hyphal-specific manner

To investigate whether Exo84-YFP is phosphorylated, we looked for a band shift on one-dimensional (1D) PAGE gels before and after λ-phosphatase treatment, using a monoclonal antibody to YFP in a Western blot experiment. Figure 3A shows that Exo84-YFP extracted from exponentially growing yeast cells has a small band shift, whereas Exo84 extracted from hyphae has a larger band shift. To investigate in more detail the pattern of phosphorylation, we fractionated Exo84-YFP using two-dimensional (2D) gels (Figure 2B). Stationary-phase yeast cells show four weak spots, whereas exponentially growing yeast cells show three further spots. Thus Exo84 is phosphorylated under yeast growth conditions consistent with the small band shift we observed in 1D gels. To study the temporal pattern of phosphorylation in hyphal cells, we analyzed samples recovered 30, 60, and 90 min after inoculation into hyphal growth conditions. Exo84 became more phosphorylated, with the spots moving progressively toward the acidic end of the pH range (Figure 3B). By 90 min, three of the additional spots appeared to correspond to the three spots visible in growing yeast, but they are more intense in the hyphal sample. A fourth spot is clearly unique to hyphal growth. Because this is the most acidic, it corresponds to the most highly phosphorylated form of the protein.

C. albicans Exo84 contains three matches to the full Cdk target motif (T/S.P.X.R/K) at positions S256, S384, and T488. Because it is known that Hgc1, the hyphal-specific Cdk1 cyclin, is required for hyphal growth, we examined whether the hyphal-specific band shift depended on the presence of Hgc1. Figure 3, B and C, shows that this is indeed the case. Exo84-YFP extracted from a Δhgc1 strain shows only a slight increase in phosphorylation compared with the stationary phase control in 2D gels (Figure 3B). In 1D gels the band shift observed in wild-type cells was absent in the Δhgc1 strain (Figure 3C).
was expressed from the regulatable MET3 promoter to investigate its possible involvement (Bensen et al., 2005). When yeast cells were grown with the MET3 promoter repressed, cells arrested with an

**FIGURE 3:** Exo84 is phosphorylated by Cdk1. (A) Western blot using anti-GFP antisera of Exo84-YFP from yeast and hyphae as indicated. Numbers indicate time in minutes since unbudded stationary-phase cells were induced to form hyphae. λPP, λ-phosphatase–treated sample. (B) 2D gels of Exo84-YFP from lysates of wild-type (WT) exo84 mutant strains as indicated and Δhgc1 strains from WT unbridged stationary-phase cells (WT H 0 min) and from hyphae induced from WT cells after 30, 60, and 90 min and Δhgc1 after 30 and 90 min as indicated. The stationary-phase cells were also reinoculated into yeast growth medium and lysates prepared after 3 h when the cells were in exponential growth (WT Yeast). (C) Exo84 extracted from Δhgc1 cells is less phosphorylated, shown by the absence of the slower-migrating smear present in Exo84 extracted from wild-type cells. (D) Exo84 is still phosphorylated in cells depleted of Clb2 and Clb4. (E) Phosphorylation of Exo84 is inhibited by 1NM-PP1 when the strain carries the cdk1-1as allele. (F) Recombinant GST-Exo84 but not recombinant GST-Exo84-3A is phosphorylated in vitro by Cdk1. The indicated recombinant forms of GST-Exo84 or GST alone were incubated in kinase reaction buffer with immunoprecipitated Cdk1-HA. The products were analyzed on a Western blot using the anti–pS-Cdk1 antibody. (G) The in vitro kinase reaction is inhibited by 1NM PP1 when the purified kinase carries the analogue-sensitive cdk1-1as allele.

In *S. cerevisiae* Cdk1 partnered by Clb2 phosphorylates Exo84 to arrest cell growth during mitosis (Luo et al., 2013). Because deletion of Clb2 is lethal in *C. albicans*, we used a strain in which Clb2
but not the presence of 1NM-PP1, Exo84 is phosphorylated in vitro, that it is not fully active. Nevertheless it is clear that in the absence lower activity in vitro even in the absence of inhibitor, suggesting allele. As we found previously (Bishop experiment using purified Cdk1-as1 carrying the analogue-sensitive kinase was Cdk1 and not a copurifying kinase, we repeated the ex 4-3A) was not a substrate (Figure 3F). To demonstrate that the active kinase is Cdk1 purified from C. albicans 3-CLB4 strain (Bensen et al., 2010), this kinase shows no change in Exo84 phosphorylation (Figure 3D). Similarly, we found no change in phosphorylation of Exo84-YFP when CLB4 expression was repressed in a MET3-CLB4 strain (Bensen et al., 2005; Figure 3D). Thus phosphorylation of Exo84 is dependent on Cdk1 partnered by Hgc1.

To further confirm the requirement for Cdk1, we examined whether the band shift was present when a strain expressing only the analogue-sensitive Cdk1-as1 allele was inhibited by the presence of 1NM-PP1. As shown in Figure 3E, the hyphal-specific band shift disappeared in the presence of 1NM-PP1. To investigate whether Cdk1 directly phosphorylates Exo84, we carried out an in vitro kinase assay with recombinant GST-Exo84 as the substrate and Cdk1 purified from C. albicans hyphae. To monitor the reaction, we used an antibody specific to phosphorylated serine in the full motif recognized by cyclin-dependent kinases (S.P.X.R/K; hereafter referred to as anti-pS-Cdk). Recombinant wild-type Exo84 was phosphorylated by purified Cdk1, but recombinant GST-Exo84 in which the three Cdk target sites had been substituted by alanine (Exo84-3A) was not a substrate (Figure 3F). To demonstrate that the active kinase was Cdk1 and not a copurifying kinase, we repeated the experiment using purified Cdk1-as1 carrying the analogue-sensitive allele. As we found previously (Bishop et al., 2010), this kinase shows lower activity in vitro even in the absence of inhibitor, suggesting that it is not fully active. Nevertheless it is clear that in the absence but not the presence of 1NM-PP1, Exo84 is phosphorylated in vitro, demonstrating that the active kinase is Cdk1 (Figure 3G).

We used the anti-pS-Cdk antibody to investigate the pattern of Cdk1-specific phosphorylation of Exo84 in vivo (Figure 4). Exo84 extracted from wild-type cells reacted strongly with this antibody from 30 min after hyphal induction onward. Exo84 extracted from Δhgc1 cells showed strongly reduced phosphorylation, consistent with the conclusion that Cdk1-Hgc1 is targeting Exo84. To confirm that the three full Cdk1 targets are involved, we constructed mutants in which these sites were mutated (described in more detail later). The strain in which all of the three Cdk1 target sites were mutated to alanine (Exo84-3A) was barely viable, producing slow-growing yeast cells that were greatly enlarged and failed to show any polarized growth in response to hyphal-inducing cues. We did not characterize this strain further. The strain in which both S256 and S384 were mutated to alanine (Exo84-2A) showed significant hyphal defects, as we document later. In a Western blot using the anti-pS-Cdk antibody, Exo4-2A showed significantly less phosphorylation than the wild type (Figure 4). We also used 2D gels to examine the pattern of phosphorylation during hyphal growth in Exo84-2A and two further mutants: one with a nonphosphorylatable alanine substitution at residue T488 (Exo84-T488A), and Exo84-3E, where the three putative phospho-acceptor sites had been mutated to phosphomimetic glutamate. In these 2D gels, in both Exo84-2A and Exo84-T488A, proteins showed reduced phosphorylation, consistent with these residues being the target of Cdk1-Hgc1 (Figure 3B). However, it was not possible to associate any particular residue with the discrete spots observed in the 2D gel of wild-type Exo84 extracted from hyphal-grown cells. Exo84-3E showed a shift toward the acidic end of the pH range, consistent with the increased negative charge of the glutamate residues.

Phosphorylation of Exo84 is required for the highly polarized growth of hyphae

To investigate the physiological role of Exo84 phosphorylation, we first confirmed that Exo84 is essential in C. albicans. In the diploid genome of C. albicans the upstream sequence of Exo84 is different in the two alleles. A retrotransposon of the Gypsy group is inserted upstream of the start of the EXO84 ORF has no major phenotypic effect; thus mutating the upstream sequence of Exo84-as1 allele was inhibited by the presence of 1NM-PP1. As shown in Figure 3E, the hyphal-specific band shift disappeared in the presence of 1NM-PP1. To investigate whether Cdk1 directly phosphorylates Exo84, we carried out an in vitro kinase assay with recombinant GST-Exo84 as the substrate and Cdk1 purified from C. albicans hyphae. To monitor the reaction, we used an antibody specific to phosphorylated serine in the full motif recognized by cyclin-dependent kinases (S.P.X.R/K; hereafter referred to as anti-pS-Cdk). Recombinant wild-type Exo84 was phosphorylated by purified Cdk1, but recombinant GST-Exo84 in which the three Cdk target sites had been substituted by alanine (Exo84-3A) was not a substrate (Figure 3F). To demonstrate that the active kinase was Cdk1 and not a copurifying kinase, we repeated the experiment using purified Cdk1-as1 carrying the analogue-sensitive allele. As we found previously (Bishop et al., 2010), this kinase shows lower activity in vitro even in the absence of inhibitor, suggesting that it is not fully active. Nevertheless it is clear that in the absence but not the presence of 1NM-PP1, Exo84 is phosphorylated in vitro, demonstrating that the active kinase is Cdk1 (Figure 3G).

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We constructed strains in which the only copy of Exo84 carried nonphosphorylatable alanine substitutions at each of the putative Cdk1 phosphorylation sites (Exo84-S256A, Exo84-S384A, Exo84-T488A). We combined the mutations to make double (Exo84-S256A S384A) and triple (Exo84-S256A S384A T488A) mutants, hereafter called Exo84-2A and Exo84-3A, respectively. We also constructed phosphomimetic substitutions at the T488 site (Exo84-T488E) and at all three putative sites (Exo84-S256E S384E T488E), hereafter called Exo84-3E. All of these strains were constructed with C-terminal YFP fusions. Fusing the remaining copy of EXO84 in the heterozygous strain to YFP had no phenotypic consequences; thus the Exo84-YFP is a functional protein. We carried out Western blots using an anti-GFP antibody, which confirmed that each of these proteins was present in amount comparable to the wild-type level (unpublished data).

As described earlier, the strain expressing only Exo84-3A was severely affected, and we did not characterize this strain further. The Exo84-2A strain showed significant morphological abnormities in the hyphal form compared with wild type (Figure 5, A and B). Quantitation showed that there was a significant increase in the number of pseudohyphal cells compared with wild type (Figure 5F), and in addition some hyphae showed a phenotype similar to the Δhgc1 phenotype, that is, a short, normal-appearing germ tube, which quickly reverted to less polarized growth resembling pseudohyphae. We refer to these cells as “other” since they resemble neither hyphae nor pseudohyphae. Although a significant proportion of Exo84-2A cells made hyphae that were morphologically normal, these hyphae were extended at approximately half the rate of wild-type hyphae (Figure 5G). Thus phosphorylation of Exo84 is necessary to allow the highly polarized growth characteristic of hyphae.

The strain expressing only the EXO84 T488A allele was similarly affected, showing marked reduction in the proportion of cells showing normal hyphal morphology (Figure 5, D and F). Again, where cells formed morphologically normal hyphae were extended at approximately half the rate of wild-type hyphae (Figure 5G). Thus phosphorylation of Exo84 is necessary to allow the highly polarized growth characteristic of hyphae. The strain expressing only the EXO84 T488A allele was similarly affected, showing marked reduction in the proportion of cells showing normal hyphal morphology (Figure 5, D and F). Again, where cells formed morphologically normal hyphae were extended at approximately half the rate of wild-type hyphae (Figure 5G). Thus phosphorylation of Exo84 is necessary to allow the highly polarized growth characteristic of hyphae.
hyphae, these hyphae extended at a markedly slower rate than the wild-type cells (Figure 5G). The strain carrying the EXO84-S384A allele also showed a reduction in the proportion of hyphal cells, and the hyphae that did form extended more slowly. However, this reduction was less extreme than for the EXO84-2A and EXO84-T488A strains (unpublished data). The EXO84-S526A allele did not have a significant effect on cell morphology or hyphal extension rate (unpublished data).

The fusion of Exo84 to YFP in the wild-type and mutant proteins allowed us to investigate the localization of Exo84. As we described previously, Exo84-YFP localizes to a surface crescent in wild-type hyphae (Figure 5, A and inset). Exo84 is also present at the septum site during cytokinesis to promote the polarized growth necessary for the formation of secondary cell walls (Figure 5A, barbed arrows). In the Exo84-2A mutant cells, Exo84-2A-YFP localized to a crescent at the tip of hyphal cells (Figure 5, B and inset). However, in cells growing in the pseudohyphal phenotype, Exo84-2A-YFP relocated to the septum late in the cell cycle and was no longer present at the tip (Figure 5B, barbed arrow), consistent with our previous report that polarity components relocate from the tip of pseudohyphal cells late in the cell cycle (Crampin et al., 2005). Exo84-T488A-YFP also localized to the tip of hyphal cells (Figure 5, C and inset) and, like Exo84-2A-YFP, was present at the tip or septum of pseudohyphal cells but not at both simultaneously. Figure 5, C and E, illustrates a phenomenon that we also occasionally observed in the Exo84-2A mutants. As can be seen in the region enlarged in Figure 5E, Exo84 becomes ectopically located with diffuse localization at the tip and an incomplete ring at the septum. Calcofluor white staining reveals that this ectopic localization is associated with excess chitin deposition. In the yeast form both the Exo84-2A and the Exo84-T488A strains showed an increase in mother cell volume compared with wild-type yeast cells (Supplemental Figure S4). However, the axial ratio (major/minor axis) was reduced, showing that cells were rounder than wild-type cells.

In hyphal-promoting growth conditions, cells expressing the phosphomimetic Exo84-3E allele were more similar to wild-type cells than cells expressing nonphosphorylatable alleles (Figure 5D). There was a smaller reduction in the proportion of cells that formed true hyphae (Figure 5F) and a smaller reduction in the rate of hyphal extension (Figure 5G). Exo84-3E yeast cells showed an increase in cell volume compared with wild-type yeast cells, but the axial ratio was not significantly different, showing that cell shape was not altered (Supplemental Figure S4). Of interest, the Exo84-T488E allele showed a decrease in cell volume and an increased axial ratio, suggesting a longer or more pronounced phase of polarized growth during their cell cycles (Supplemental Figure S4).

Whereas nonphosphorylatable alanine substitutions at the Cdk1 target sites impair polarized growth and hyphal morphology, phosphomimetic substitutions have only a mild effect on the hyphal phenotype. This is consistent with the hypothesis that phosphorylation of Exo84 is necessary for the extreme polarized growth of hyphae. If one of the roles of Hgc1-Cdk1 is to phosphorylate Exo84, then it is possible that the Exo84-3E allele may alleviate some or all of the phenotypic effects of the Δhgc1 allele. To test this idea, we constructed a Δhgc1 EXO84-3E strain. The EXO84-3E allele did not rescue the phenotype of the Δhgc1 allele. However, there was an increase in the proportion of hyphal cells from 1.2 to 17% (Figure 5F). Because other functions of Hgc1-Cdc28 have been documented, it is not surprising that the EXO84-3E allele does not fully rescue the effects of a Δhgc1 mutation. One such target of Hgc1-Cdk1 is Rga2, which acts as a negative regulator of hyphal growth that is relieved upon phosphorylation (Zheng et al., 2007). We previously showed that although deletion of Rga2 does not induce constitutive hyphal growth, hyphal growth is initiated in Δrga2 mutants under conditions that would normally induce pseudohyphal formation (Court and Sudbery, 2007). We thus investigated whether the Exo84-3E mutant also increased the proportion hyphae when cells were grown at 35°C and pH 6.0, which would normally induce pseudohyphal growth (Figure 6). Indeed, under these conditions the proportion of hyphae increased from 15% in wild-type cells to 35% in Exo84-3E mutants. In contrast, hyphal growth was absent (Exo84-T488A) or nearly absent (Exo84-2A) in the nonphosphorylatable mutant alleles (Figure 6). Taken together, these experiments provide further support for the hypothesis that Hgc1-Cdk1 targets Exo84 at the three full consensus Cdk1 target sites to promote hyphal growth, although it is clear that Cdk1 also has other targets that must be phosphorylated for the full hyphal phenotype.

When hyphae are able to form, phosphorylation of Exo84 is not required for its localization or exocyst assembly

Exo84 is continuously present as a surface crescent at the tip of hyphae, and during cytokinesis it is simultaneously present at the site of septum formation and the hyphal tip (Figures 5A). The morphological defects resulting from the mutant alleles described earlier were observed in cells growing on rich yeast extract/peptone/glucose (YEPD) medium. We found that the morphological defects of the exo84-2A and exo84-T488A alleles were less pronounced on minimal medium, so that a higher proportion had a hyphal morphology (Figure 7A). Nevertheless, these hyphae extended at a slower rate than wild-type hyphae, showing that the nonphosphorylatable alleles still reduced polarized growth. We took advantage of this observation to investigate whether the mutations affected the location of Exo84 to sites of septum formation during hyphal growth. We followed Exo84-YFP localization over time in wild-type cells and strains carrying each of the nonphosphorylatable and phosphomimetic alleles fused to YFP. Figure 7B shows the percentage of cells displaying Exo84 in the septum plotted against time after hyphal induction. In wild-type cells a peak of septum-located Exo84 occurs at 115 min. Cumulatively, Exo84 was observed at the septum in 70% of wild-type hyphae by 150 min (Figure 7C). Mutant cells that formed hyphae also showed localization of Exo84 to the septum (Figure 7A). However, it is observed at a later time in the septum compared with wild-type cells, and the cumulative proportion of cells that displayed septum-located Exo84 is much lower in each of the mutants (Figure 7C). However, when Exo84 is present at the septum, the total length of the hypha and the distance from the bud neck to the septum is similar in the wild type and mutants (Figure 7D). This suggests that the delay in Exo84 location to the septum reflects an increased time to reach a critical hyphal length at which cytokinesis is initiated due to the lower extension rate of the hyphae. Thus in these hyphal cells the presence or absence of phosphorylation does not affect Exo84 localization but does affect the rate of polarized growth.

In S. cerevisiae, phosphorylation of Exo84 disrupts the assembly of the exocyst complex (Luo et al., 2013). We showed (Supplemental Figure S2) that Exo70 remains at the bud tip throughout mitosis. To further investigate the effect of Exo84 phosphorylation on exocyst interactions, we examined whether phosphorylation of Exo84 affected its interaction with Sec10, as this interaction is disrupted in S. cerevisiae by Exo84 phosphorylation. We constructed Exo84-GFP Sec10-HA strains in which Exo84 was wild type or carried respectively the Exo84-T488A, Exo84-2A, and 3E mutations. Using these strains, we carried out reciprocal immunoprecipitation experiments. In each case the mutations did not affect the interaction of Exo84 and Sec10 (Figure 8, A and B).
Exo84 affects the binding. To do this, we used a PIP strip and recombinant GST-Exo84 and GST-Exo84-3E (Figure 9A). Wild-type GST-Exo84 appeared to have the greatest affinity for phosphatidylserine (PS), with a much weaker affinity for phosphatic acid (Figure 9, A and B). Because this protein was expressed in Escherichia coli, it is not phosphorylated, and so this reports the affinity of the nonphosphorylated form of the protein. Of interest, the GST-Exo84-3E protein appeared to have a considerably reduced affinity for PS, consistent with the negative charge on PS and the increased negative charge on Exo84 resulting from the phosphorylation (Figure 9, A and B). Note that 8 μg of recombinant Exo84-3E but only 2 μg of recombinant Exo84 were loaded onto each spot in the PIP strip; thus the reduced affinity of Exo84 for PS is likely to be considerably greater than the 2.5-fold reduction in Figure 9A.

PS is synthesized from cytidyldiphosphate-diacylglycerol and serine by the action of phosphatidylserine synthase, encoded by the CHO1 gene (Chen et al., 2010). In C. albicans, Δcho1 mutants are viable but have cell wall defects and fail to show filamentous growth on Spider medium. In addition, they are auxotrophic for ethanolamine, as phosphatidylethanolamine is synthesized from PS. To investigate further whether PS plays a role in Exo84 localization, we constructed a Δcho1 Exo84-YFP strain and investigated its phenotype. We used defined medium supplemented with phosphatidylethanolamine, since we found that YEPD contained sufficient PS to supplement the PS auxotrophy. On defined medium this strain grew slowly. In both wild-type and Δcho1 yeast cells Exo84-YFP localized to the septum and the tip of small buds (Figures 7A and 9C). During hyphal growth in Δcho1 cells (induced on Lee’s medium; Lee et al., 1975), Exo84 also showed a disturbed pattern of localization, so that although it was present at the tip, it was also present in ectopic sites in the mother cell and within the germ tube (Figure 9C). As a control, we also constructed a Δcho1 strain expressing Sec3-YFP, which during both yeast and hyphae growth did not show the ectopic localization. (Figure 9C). Thus ectopic localization of Exo84 in the absence of PS is specific to Exo84 is not caused by a general failure of exocyst localization.

Although these observations suggest a role for PS in the localization of Exo84, the role of this phosphorylation is not straightforward because phosphorylation lowers the affinity of Exo84 for PS. It may be that the phosphorylation is important to dissociate Exo84 from membranes to allow it to be recycled to the growing tip in a manner reminiscent of recent models of the role of endocytosis in restricting

Phosphorylation of Exo84 may affect phospholipid binding
Exo84 contains a PH domain that promotes interactions between proteins and phospholipids (Figure 1A). The S384 site is located within this domain, whereas protein threading suggests that the important S256 and T488 sites are on either side of this domain in the folded protein (Figure 1A). We therefore investigated whether Exo84 binds phospholipids, and if so, whether phosphorylation

FIGURE 6: The phosphomimetic EXO84-3E allele promotes hyphal growth under pseudohyphal growth conditions. Stationary-phase cells were inoculated into YEPD, pH 6.0, and cultured at 35°C. Morphology was classified as described in the legend to Figure 5. A minimum of 150 cells was classified for each strain.

Exo84 phosphorylation by Cdk1
During the growth of a bud in S. cerevisiae or the yeast phase of C. albicans, growth is polarized for a short period as the bud emerges but later becomes isotropic around the whole of the bud perimeter. The period of polarized growth relative to isotropic growth must be relatively short because, as we record here, the axial ratio of the prolate spheroid C. albicans cells is 1.5. During cytokinesis, polarized growth relocates to the site of septum formation. However, it was recently shown in S. cerevisiae that fusion of secretory vesicles is blocked during metaphase, and cell surface expansion ceases (Luo et al., 2013). C. albicans hyphae show a radically different pattern of growth than the yeast phase. Polarized growth is focused to a narrow area at the hyphal tip and is continuous throughout the whole cell cycle. It does not show the polarized-to-isotropic switch, and we show here that it does not cease at any point during mitosis or cytokinesis. Despite this radically different growth pattern, studies on the mechanism of polarized growth in C. albicans suggest that the same components are used as elaborated in the S. cerevisiae budding yeast model (for review see Sudbery, 2011). Clearly, however, the machinery must be modified to produce the radically different outcome of hyphal growth.

Here we uncovered one of the key differences. In C. albicans the exocyst component Exo84 is phosphorylated by Cdk1 as it is in S. cerevisiae. However, instead of causing the disassembly of the exocyst complex and halting the fusion of secretory vesicles necessary for cell surface expansion, phosphorylation of Exo84 is necessary for the efficient polarized growth of hyphae.

Substitutions of nonphosphorylatable residues at the consensus Cdk1 phosphorylation sites cause morphological abnormalities in many cells growing on rich medium and reduces hyphal extension rates on all media. The morphological abnormalities include an localization of Cdc42 to sites of polarized growth. To address this possibility, we examined the recycling kinetics of Exo84-2A and Exo84-3E using FRAP. To avoid the application of extensive bleaching corrections during postbleach imaging, we used an experimental design we developed previously in which we measured the fraction of fluorescence recovered in 30 s (Jones and Sudbery, 2010). Exo84-2A-YFP recovered only 35% of the prebleach fluorescence in 30 s, whereas wild-type Exo84-YFP recovered 53%, a difference that was statistically significant (Figure 10). Exo84-3E recovered 60% of prebleach fluorescence. Although this was more than the wild-type protein, the difference was not significant. Thus phosphorylation of Exo84 is necessary for the rapid exchange of Exo84 at the cell surface, consistent with the idea that phosphorylation promotes Exo84 recycling by promoting its dissociation from PS.
We also observed that Exo84 is phosphorylated during yeast-phase growth, although this was not as extensive as hyphal growth. Exo84 phosphorylation is clearly important for yeast growth, as shown by the major effect of the Exo84-3A allele in the yeast-phase phenotype; moreover, the mother cells of Exo84-3A, Exo84-2A, and Exo84-T488A strains were rounder than the wild type. This suggests that Exo84 phosphorylation is required for the initial polarized growth of small yeast buds. Intriguingly, it was previously reported that S. cerevisiae cells arrested at Start by cdc28-4 and cdc7-1 alleles at 37°C show increased affinity of Exo84 for other components of the exocyst (Luo et al., 2013). In addition, Exo84 shows a weak reaction to the antibody recognizing phosphorylated Cdk1 target sites in cells arrested at start by α-factor. Such cells form mating projections that have many similarities to hyphae (Chapa-Y-Lazo et al., 2011). Finally, the Cln2-Cdk1 kinase phosphorylates recombinant Exo84 in vitro (Luo et al., 2013). Taken together, these observations suggest that phosphorylation of Exo84 may also play a role in polarized growth in S. cerevisiae.

What is the role of Exo84 phosphorylation? One of the Cdk sites is within the PH domain, and the other two sites are located on either side of the PH domain in the 3D model of the protein reconstructed by threading analysis (Figure 1A). PH domains bind phospholipids in membranes, and we show here that recombinant GST-Exo84, which is not phosphorylated, has an affinity for PS and that this is reduced in the Exo84-3E, which mimics a fully phosphorylated protein. Thus phosphorylation may regulate the affinity of Exo84 for this phospholipid. A precedent for the role of PS in polarized growth is the demonstration that in S. cerevisiae it is localized to sites of polarized growth and facilitates recruitment of Cdc42-GTP to these sites. We showed that in Δcho1 cells, which are unable to synthesize PS, and that this is reduced in the Exo83-3E, which mimics a fully phosphorylated protein. Thus phosphorylation may regulate the affinity of Exo84 for this phospholipid. A precedent for the role of PS in polarized growth is the demonstration that in S. cerevisiae it is localized to sites of polarized growth and facilitates recruitment of Cdc42-GTP to these sites. We showed that in Δcho1 cells, which are unable to synthesize PS, Exo84 localized ectopically to punctate patches at the cell surface in both yeast and hyphae. Nevertheless, in these cells Exo84 also localizes to sites of polarized growth such as the tip of small buds and the septum in yeast, and the hyphal tip and septum in hyphae. Thus loss of PS lowers the efficiency of, but does not completely prevent, Exo84 localization. In S. cerevisiae, Sec3 is believed to provide a landmark for exocyst assembly. In C. albicans, ΔSec3 cells fail to maintain polarized growth after the formation of the first septum, a phenotype that depends on the presence of Cdc11 in the septin ring (Li et al., 2007). In C. albicans Δcho1 cells, Sec3 localizes normally to the septum, showing that the inefficient localization of Exo84 in Δcho1 cells is specific and not due to general disorganization of the polarity machinery. A possible role for the phosphorylation is that it promotes increased proportion of cells that grow as pseudohyphae, as well as cells that resemble the Δhgc1 phenotype, where cells evaginate germ tubes that appear normal but quickly revert to the less-polarized growth typical of pseudohyphae. These abnormalities are not shown by cells carrying phosphomimetic substitutions at the same sites. In particular, whereas the Exo84-3A mutant is severely affected and in many ways resembles the phenotype of cells depleted of Exo84, the Exo84-3E strain shows only minor abnormalities.
reduction in its affinity for PS, whereas ScExo84 contains a phosphorylation site in the C-terminal interaction domain, explaining the loss of interaction with other exocyst subunits. Thus, whereas EXO84 is present in the genomes of both species, it has evolved so that its phosphorylation adapts each organism to its particular pattern of growth.

MATERIALS AND METHODS

Media and growth conditions

YEPD consists of 2% glucose, 2% Bacto peptone (Difco), and 1% Bacto yeast extract (Difco) plus 80 mg/l uridine. SD medium consists of 0.67% (wt/vol) yeast nitrogen base (Difco), 2% (wt/vol) glucose, 80 mg/l uridine or 40 mg/l histidine, and arginine. Hyphae were induced from un-budded stationary-phase yeast cells as described previously. The Δcho1 mutant was grown in synthetic defined (SD) medium containing 1 mM ethanolamine, and hyphal growth was induced in Lee’s medium (Lee et al., 1975). Shutdown of pMET3-CLB2 experiments was carried out in SD supplemented with 10 mM methionine and 2.5 mM cysteine.

Strain and plasmid constructions

Strains constructed are listed in Supplemental Table S1, and the oligonucleotides used are listed in Supplemental Table S2. All strains were derived from BWP17 (Wilson et al., 1999). Gene deletions and C-terminal YFP fusions and hemagglutinin (HA) fusions were performed as previously described (Gola et al., 2003; Schaub et al., 2006; Lavoie et al., 2008; Walther and Wendland, 2008). All strains were checked for correct genome integration by PCR. Correct expression of protein fusion strains was also verified by Western blot using antibodies to the fusion protein or epitope (unpublished data).

Western blots

Western blots were carried out as described previously (Wightman et al., 2004).

2D gel electrophoresis

Exo84-YFP was immunoprecipitated as previously described and then eluted from beads with hydration buffer (8 mM urea, 2 M thiourea, 4% 3-[3-cholamidopropyl]dimethylammonio]-1-propanesulfonate, 1 mM dithiothreitol [DTT], 0.5% IPG buffer, pH 3–10, and 1.2% DeStreak Reagent; GE Healthcare). The immunoprecipitated protein was resolved on 2D gels using 7-cm Immobiline Drystrip, pH 3–10, and the Ettan IPGphor 3 Isoelectric Focusing System (GE Healthcare).
FIGURE 10: Nonphosphorylatable substitutions reduce Exo84-YFP recovery after photobleaching. The indicated strains were grown as hyphae on agar pads in a DeltaVision microscope (TA, Exo84-EXO84 T488A). The tips were bleached using a laser. Images were recorded prebleach, postbleach, and postbleach plus 30 s. Exo84-YFP fluorescence was quantified as the fractional area of the tip above background fluorescence intensity. Recovery was defined as (Ib - Io)/(Ib - Io) x 100%, where Ib, Io, and I0 are the values of Exo84-YFP fluorescence prebleach, postbleach, and 30-s postbleach, respectively.

Exocyst complex coimmunoprecipitation

Cells were broken in IP-Lysis buffer (5% glycerol, 50 mM Tris-HCl, pH 7.5, 150 mM KCl, 2 mM MgCl2, 0.5 mM ethylene glycol tetraacetic acid, 1 mM DTT, and protease inhibitor tablet [Roche]). Immunoprecipitation was done as described, using anti-GFP or anti-HA antibodies.

In vitro kinase assays

Cdc28-HA or Cdc28as-HFP was immunoprecipitated in hyphal lysates using anti-HA (12CA5) monoclonal antibody. Kinase assays were carried out as previously described (Bishop et al., 2010). The anti–pS-Cdk antibody was used to detect Cdk1-specific phosphorylation.

Phospholipid-binding assay

Recombinant glutathione S-transferase (GST)–Exo84, GST-Exo84-3A, and GST-Exo84-3E were expressed in E. coli and purified with glutathione Sepharose 4b (GE Healthcare) according to the manufacturer’s instructions. For the strips shown in Figure 9C, 8 μg of purified GST-Exo84-3E and 2 μg of purified GST-Exo84 protein were incubated with a PIP Strip (Echelon Biosciences, Salt Lake City, UT) previously blocked with 3% bovine serum albumin. Protein binding was visualized using anti-GST antibody (Santa Cruz Biotechnology, Dallas, TX).

Microscopy

Live-cell imaging was carried out on cells growing on agar pads using a DeltaVision RT microscope (Applied Precision Instruments, Seattle, WA) as described previously (Bishop et al., 2010). The intensity of the Mic1-YFP signal in the Spitzenkörper shown in Figure 2 was measured using the plot Z-axis profile tool in the FIJI distribution of ImageJ (http://fiji.sc/Fiji) after a region of interest (ROI) had been drawn around the area occupied by the Spitzenkörper in frames 43–58 of Supplemental Movie S2. The data are shown with the background measured within the cell subtracted. The images shown in Figure 5 were obtained from cells grown in YEPD liquid culture 120 min after hyphal induction and stained in phosphate-buffered saline (PBS) buffer with Calcofluor white (Fluorescent Brightener 28; Sigma-Aldrich) at a concentration of 1 μg/ml. To quantify cell dimensions and morphology shown in Figure 5, hyphal cells were grown for 120 min in YEPD liquid culture as described. After harvesting, the cells were fixed with 2% formaldehyde and then treated with pepsin to dissociate clumped hyphae as previously described (Sudbery, 2001). Cell dimensions were measured using FIJI aided by an in-house custom script that recorded cell dimensions and user-defined definitions of morphological state (hyphae, pseudohyphae, or other). Calcofluor white (Fluorescent Brightener 28) was added to liquid grown cells at a concentration of 1 μg/ml. FRAP experiments were carried out using a DeltaVision Microscope with an attached 532-nm laser module for bleaching. Images were analyzed in FIJI as follows. For each image a threshold was set to exclude the background. An ROI was drawn around the crescent of Exo84 in the prebleach image. The fractional area above the threshold in prebleach, postbleach, and postbleach plus 30-s images was then recorded. Recovery was defined as (Ib – Io)/(Ib – Io) x 100%, where Ib, Io, and I0 are the values of Exo84-YFP intensity prebleach, postbleach, and 30-s postbleach, respectively.

Protein threading

The Protein Homology/Analogy Recognition Engine, version 2.0 (Phyre2), in its intensive modeling mode was used to obtain the Exo84 structure (Kelley and Sternberg, 2009). Analysis, formatted, and final movies and images were created using PyMOL (PyMOL Molecular Graphics System, version 1.2.3pre; Schrödinger).

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