Phosphatidylserine synthesis is essential for viability of the human fungal pathogen Cryptococcus neoformans

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Edited by F. Peter Guengerich

Phospholipids are an integral part of the cellular membrane structure and can be produced by a de novo biosynthetic pathway and, alternatively, by the Kennedy pathway. Studies in several yeast species have shown that the phospholipid phosphatidylserine (PS) is synthesized from CDP-diacylglycerol and serine, a route that is different from its synthesis in mammalian cells, involving a base-exchange reaction from preexisting phospholipids. Fungal-specific PS synthesis has been shown to play an important role in fungal virulence and has been proposed as an attractive drug target. However, PS synthase, which catalyzes this reaction, has not been studied in the human fungal pathogen Cryptococcus neoformans. Here, we identified and characterized the PS synthase homolog (Cn Cho1) in this fungus. Heterologous expression of Cn CHO1 in a Saccharomyces cerevisiae cho1Δ mutant rescued the mutant’s growth defect in the absence of ethanolamine supplementation. Moreover, an Sc cho1Δ mutant expressing Cn CHO1 had PS synthase activity, confirming that the Cn CHO1 encodes PS synthase. We also found that PS synthase in C. neoformans is localized to the endoplasmic reticulum and that it is essential for mitochondrial function and cell viability. Of note, its deficiency could not be complemented by ethanolamine or choline supplementation for the synthesis of phosphatidylethanolamine (PE) or phosphatidylcholine (PC) via the Kennedy pathway. These findings improve our understanding of phospholipid synthesis in a pathogenic fungus and indicate that PS synthase may be a useful target for antifungal drugs.

Cryptococcus neoformans is a human fungal pathogen and the leading cause of life-threatening fungal meningoencephalitis, especially in immunocompromised individuals (1, 2). It is an environmental organism that infects humans via the respiratory tract to cause pulmonary cryptococcosis, followed by homogeneous dissemination to the central nervous system to cause meningitis (3). Despite its clinical significance, treatment options for invasive cryptococcosis are extremely limited. Currently, an acute infection is commonly treated with amphotericin B or azoles in combination with 5-flucytosine (4). However, both treatment courses have serious drawbacks: amphotericin B produces toxic side effects, whereas triazoles are fungistatic, which can promote the emergence of drug resistance. The main therapeutic challenge in developing new antifungal agents is that both fungi and their mammalian hosts are eukaryotes and therefore contain similar cellular machinery. Hence, identification and characterization of fungal-specific enzymes that are important for fungal survival, growth, or virulence could identify valuable potential drug targets.

Phospholipids are an integral part of eukaryotic cell membrane and are asymmetrically distributed across the bilayer. In general, phosphatidylserine (PS)2 and phosphatidylethanolamine (PE) are mainly restricted to the cytoplasmic leaflet of the membrane, whereas phosphatidylcholine (PC) and sphingolipids are mainly concentrated in the exocytosomal leaflet (5, 6). The lipid asymmetry creates two distinct membrane surfaces with very different adhesive properties. Among them, PS is the most abundant negatively charged phospholipid in eukaryotic membranes. Its covalent attachment of serine to the phosphate gives PS a net negative charge on the head group (7, 8). A high PS concentration in the cytoplasmic leaflet of the plasma membrane in mammals is essential for targeting and function of several intracellular signaling proteins and activation of specific kinases, such as protein kinase C (6, 9). Translocation of PS to the exocytosomal leaflet of the membrane could be detrimental to cell viability, because macrophages recognize cell-surface PS as a phagocytic signal to engulf cells by binding to the exposed PS (8, 10, 11). In mammals and some parasites (e.g. Trypanosoma brucei), PS is generated by exchanging serine for choline in PC and for ethanolamine in PE by the activity of PS synthases PSS1 and PSS2, respectively (12, 13). In yeast, however, PS is synthesized from CDP-diacylglycerol (CDP-DAG)

2 The abbreviations used are: PS, phosphatidylserine; PE, phosphatidylethanolamine; PC, phosphatidylcholine; DAG, diacylglycerol; SC, synthetic complete; H2DCFDA, dichlorodihydrofluorescein diacetate; DAPI, 4′,6-diamidino-2-phenylindole; ROS, reactive oxygen species; 1D-TLC, one-dimensional thin-layer chromatography; ER, endoplasmic reticulum; qRT, quantitative RT; BCS, bathocuproinedisulfonic acid.
and serine by the fungal-specific PS synthase enzyme Cho1 (14–16).

Thus far, the role of Cho1 has been investigated in Saccharomyces cerevisiae and Candida albicans. The S. cerevisiae cho1 mutant (Sc cho1Δ) is viable, but because it is unable to synthesize PS, which serves as a substrate for the de novo synthesis of PE and PC, the Sc cho1Δ mutant can only grow if ethanolamine or choline is exogenously supplied, enabling PE and PC synthesis via the Kennedy pathway (Fig. 1) (17). The Sc CHO1 gene expression is negatively regulated by inositol and choline availability (18). The C. albicans cho1Δ (Ca cho1Δ) mutant can only grow when supplemented with ethanolamine (15), but it cannot be rescued by choline on solid medium and displays poor growth when supplemented with choline in liquid medium. In addition, the Ca cho1Δ mutant is avirulent in a mouse model of systemic candidiasis, suggesting that the PS synthase homolog in C. albicans is necessary for virulence and therefore represents a potential new drug target (15). However, the function and importance of PS synthase in C. neoformans development and pathogenesis have not been analyzed yet.

Our previous study revealed that C. neoformans lipid flipase, which translocates certain phospholipids (PS and PE) from the exocellular leaflet to the cytoplasmic leaflet to maintain the asymmetrical distribution of phospholipids, is essential for virulence in a murine model of cryptococcosis. It is also required for the inherent resistance of this fungus to echinocandin drugs. Mutants of the lipid flipase regulatory subunit Cdc50 showed accumulation of PS on the outer layer of the plasma membrane, were hypersensitive to macropage killing, and showed reduced virulence (19). We hypothesized that loss of Cdc50 prevents PS translocation, leading to increased PS exposure on the cell surface, which may promote macrophage recognition and killing of such cells, because PS accumulation on the cell surface has been shown to serve as a phagocytic signal for macrophages (20). The overall reduced phospholipid level in the cdc50Δ mutant was also observed in a separate study (21). Therefore, we were interested in further exploring the role of PS in fungal development and virulence.

In this study, we characterized the only Cho1 homolog in C. neoformans (Cn Cho1) and confirmed that it functions as a PS synthase. We were unable to generate a deletion mutant of Cn CHO1, revealing that Cn Cho1 is essential for C. neoformans viability, despite the presence of an active Kennedy pathway in this fungus. We further investigated the mechanism of Cn Cho1 essentiality and revealed that PS is likely an essential phospholipid for this fungus. The essentiality of the fungal-specific PS synthase in C. neoformans warrants further investigation of this enzyme as a potential drug target.

**Results**

**Cho1 homolog in C. neoformans is a PS synthase**

Our previous study showed that deletion of the regulatory subunit of the lipid flipase Cdc50 leads to the accumulation of PS on the outer leaflet of the bilayer membrane in *Cryptococcus*, which appears to contribute to the loss of fungal virulence (19). Therefore, we tried to generate a mutant strain lacking PS to better understand the role of PS in fungal virulence. This led us to identify the only homolog of the *S. cerevisiae* PS synthase Cho1 protein.

The C. neoformans Cho1 (Cn Cho1) protein shares high amino acid sequence identity with the known PS synthases in *S. cerevisiae* (Sc Cho1) and *C. albicans* (Ca Cho1). All of them share the conserved sequence of CDP-diacylglycerol-serine O-phosphatidyltransferase (DGX…ARX…GX3DX3D). Protein secondary structure analysis also indicated that all three are integral membrane proteins with multiple transmembrane domains, albeit with some differences in the number and the topology of potential transmembrane regions (Fig. 2).

To confirm the localization of Cn Cho1 protein in *C. neoformans*, we generated a Cn Cho1:mCherry fusion construct and expressed this fusion protein in *C. neoformans* WT H99. Our results showed that the fluorescent signal mostly localized to the ER membrane (Fig. 3A and Fig. S1). We then purified the membrane fraction by centrifugation and treated it with either PBS, Triton X-100, or 1M NaCl to determine whether Cn Cho1 is an integral membrane protein or a membrane-binding protein (Fig. 3B). Our results clearly showed that Cn Cho1 was solubilized by Triton X-100, but not by high salt, indicating that it is an integral membrane protein.

To confirm the Cn Cho1 protein is a PS synthase, we expressed Cn CHO1 gene in a *S. cerevisiae* strain carrying a deletion of Sc CHO1 (Sc cho1Δ). As expected, the Sc cho1Δ mutant was an ethanolamine auxotroph and could not grow on SC medium lacking ethanolamine (Fig. 4A). We observed that heterologous expression of Cn CHO1 in this mutant background fully complemented its growth defect, indicating that Cn CHO1 is likely a PS synthase (Fig. 4A). We then measured the PS synthase activity in cell lysates directly using [3-3H]serine as described previously (16, 22, 23). Our results showed that the Sc cho1Δ mutant completely lacked PS synthase activity and that expressing Cn CHO1 in this mutant largely restored its PS synthase activity (Fig. 4B), supporting the conclusion that Cn Cho1 has PS synthase activity.

In *S. cerevisiae*, extracellular inositol negatively regulates Sc CHO1 expression (18). Thus, we examined Cn CHO1 gene expression in the presence of various inositol concentrations to determine whether it is also regulated by inositol. Indeed, our results indicated that Cn CHO1 transcription is repressed by a high level of inositol and induced or derepressed as the inositol
level decreases (Fig. 4C). Although we have not been able to identify C. neoformans homologs of S. cerevisiae inositol-dependent transcriptional regulators, our results indicate that the Cn CHO1 gene is regulated by inositol similarly to Sc CHO1.

**CHO1 is an essential gene in C. neoformans**

To examine the role of PS synthase in C. neoformans, we attempted to generate a deletion mutant for Cn CHO1 in the H99 background, but were unsuccessful despite the extensive effort of screening over 200 transformants (data not shown). We then deleted one allele of Cn CHO1 from a diploid C. neoformans strain AI187 (24), replacing it with a neomycin (NEO) resistance cassette (Fig. 5A). The heterozygous Cn CHO1/cho1/H9004 strain was viable and grew normally on YPD. We incubated the strain on a V8 mating medium to induce mating and sporulation, dissected over 20 spores, and patched them on YPD with or without G418. All of the tested spores grew on YPD, but none of them grew on YPD with G418, suggesting that all viable spores contain the WT Cn CHO1 allele (Fig. 5C) and that Cn Cho1 is likely essential for fungal viability. We also repeated the experiment by supplementing the medium with 1 mM ethanolamine or 1 mM choline, but we still did not obtain any viable haploid Cn cho1Δ strains (data not shown).

To better understand the potential essentiality of Cn Cho1, we used homologous recombination to replace the Cn CHO1 native promoter with an inducible CTR4 promoter (PCTR4-CHO1). The activity of the CTR4 promoter can be suppressed by the addition of CuSO4 and ascorbic acid and induced by the addition of the copper chelator bathocuproinedisulfonic acid (BCS) (25). First, we tested cell growth by adding 25 μM CuSO4 that has been shown to significantly suppress the CTR4 promoter in previous studies (25, 26). However, we did not observe a significant growth defect between WT and the PCTR4-CHO1 strain (Fig. S3). By testing the growth assay using media containing different CuSO4 concentrations, we found that when 1 mM CuSO4 was used, the strain grew very poorly (Fig. 6, B and C). As a control, WT strain treated with 1 mM CuSO4 only showed modest growth defect (Fig. 6D), which is consistent with a previous report using such a CuSO4 concentration (27).

qRT-PCR analysis of CHO1 expression under the control of the CTR4 promoter showed that although 25 μM CuSO4 significantly inhibited the gene expression, adding 1 mM CuSO4 led to even further inhibition (Fig. 6A). From these results, we concluded that although 25 μM CuSO4 can significantly reduce gene expression, the remaining low level of PS synthase activity...
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is sufficient to support fungal growth. The further gene suppression due to addition of 1 mM CuSO₄ may be sufficient to abolish the Cho1 enzyme activity, leading to growth arrest.

Figure 3. Cn Cho1 is an integral membrane protein that primarily localizes to the endoplasmic reticulum. A. Cho1 protein in C. neoformans localizes to the ER membrane in a similar manner to Cho1 from other yeasts. In this strain, the CHO1 gene is fused with an mCherry tag at its C terminus, and its expression is under the control of the C. neoformans actin promoter. ER tracker was used to visualize the ER membrane. Among over 100 cells analyzed, ~99% showed co-localization of Cho1 protein with the ER tracker. Additional data were shown in Fig. S1. Scale bar, 10 μm. B, membrane protein separation assay confirms that Cho1 is an integral membrane protein. As described under “Experimental procedures,” a total membrane fraction obtained from cells expressing Cho1 was extracted with 1× PBS buffer, 1% Triton X-100, or salt (1 M NaCl). Following ultracentrifugation, equivalent amounts of the insoluble pellet (P) and the supernatant (S) were analyzed by Western blotting using RFP-tag antibody.

Figure 4. C. neoformans Cho1 is a PS synthase, and its expression is regulated by inositol. A, expression of Cn CHO1 gene fully complements S. cerevisiae (Sc) cho1Δ mutant auxotrophic for ethanolamine (Etn). WT S. cerevisiae can grow well in the absence of ethanolamine unlike the Sc cho1Δ mutant transformed with pTH19 (empty vector), which is ethanolamine auxotrophic. Sc cho1Δ expressing Cn CHO1 restores growth in the absence of ethanolamine, indicating that C. neoformans CHO1 gene can fully complement Sc CHO1 gene function. B, enzymatic analysis of Saccharomyces strains for PS synthase activity. Cho1 enzyme activity in WT S. cerevisiae was absent in Sc cho1Δ strain. This activity was successfully restored when the Sc cho1Δ strain was transformed with Cn CHO1 gene indicating that it is a PS synthase. The assay was performed in duplicate and repeated twice. C. Cn CHO1 gene expression is regulated by inositol. CHO1 gene expression is suppressed in the presence of 1% inositol as compared with its expression in the presence of 0.1% inositol indicating that phospholipid precursor inositol negatively regulates CHO1 gene expression. The analysis of CHO1 gene expression was performed in triplicate, and the two-tail t test was used for statistical analysis (* indicates p < 0.001).

Figure 5. Cn Cho1 is an essential protein for fungal viability. A Cn CHO1/ cho1Δ heterozygous mutant was generated by deleting one Cn CHO1 allele in a diploid strain AI187 and undergoing sporulation on mating medium. Spores were isolated and grown on YPD or YPD + G418. Among the 28 spores isolated, none of them grew on YPD containing G418, indicating that they are all WT alleles. A, schematic of Cn CHO1 deletion strategy. CX numbers indicate primers used to screen the mutants. Detailed description of each primer can be found in Table S1. B, PCR results of the mutant screen. M1, mutant 1; M2, mutant 2. C, growth of progeny colonies on YPD and YPD containing G418 after 3 days of incubation.
The Kennedy pathway in *C. neoformans* is active in *S. cerevisiae*. Phospholipids such as PE and PC can be produced either by decarboxylation of PS or by utilizing ethanolamine and choline via the Kennedy pathway (Fig. 1) (17). Therefore, *S. cerevisiae cho1Δ* mutant is ethanolamine and choline auxotrophic and is still viable when supplemented with ethanolamine or choline. We investigated whether ethanolamine or choline could also rescue the growth arrest of the *PCTR4-CHO1* strain under conditions that repressed CHO1. Interestingly, we found that adding 1 mM ethanolamine or 1 mM choline did not rescue the growth of the *PCTR4-CHO1* strain under inhibition conditions (Fig. 6, B and C). One possibility for this lack of rescue is that the Kennedy pathway may be inactive in *C. neoformans*. To test this possibility, we monitored PE and PC production by using [14C]ethanolamine and [14C]choline in a 1D-TLC analysis. Our data showed that both *C. neoformans* WT H99 and *S. cerevisiae* WT BY4742 could produce PE and PC from ethanolamine and choline, respectively, indicating that the Kennedy pathway is active in *C. neoformans* (Fig. 7A).

### The Kennedy pathway in *C. neoformans* is active

In *S. cerevisiae*, phospholipids such as PE and PC can be produced either by decarboxylation of PS or by utilizing ethanolamine and choline via the Kennedy pathway (Fig. 1) (17). Therefore, *S. cerevisiae cho1Δ* mutant is ethanolamine and choline auxotrophic and is still viable when supplemented with ethanolamine or choline. We investigated whether ethanolamine or choline could also rescue the growth arrest of the *PCTR4-CHO1* strain under conditions that repressed CHO1. Interestingly, we found that adding 1 mM ethanolamine or 1 mM choline did not rescue the growth of the *PCTR4-CHO1* strain under inhibition conditions (Fig. 6, B and C). One possibility for this lack of rescue is that the Kennedy pathway may be inactive in *C. neoformans*. To test this possibility, we monitored PE and PC production by using [14C]ethanolamine and [14C]choline in a 1D-TLC analysis. Our data showed that both *C. neoformans* WT H99 and *S. cerevisiae* WT BY4742 could produce PE and PC from ethanolamine and choline, respectively, indicating that the Kennedy pathway is active in *C. neoformans* (Fig. 7A).

### PS is essential for Cryptococcus cell viability

Another possibility for *Cn CHO1* essentiality is that PS is essential for viability in *C. neoformans*. Although PE and PC can be produced by both the *de novo* PS synthetic pathway and the Kennedy pathway, PS can only be produced via PS synthase (Cho1). Therefore, we supplemented the growth medium with 1 mM lyso-PS and examined whether lyso-PS (derivative of phosphatidylserine with one fatty acid chain that rapidly incor-
Porates into the cells and is converted to PS) could rescue the growth arrest induced by CuSO₄. Indeed, we observed significant rescue of growth defect by lyso-PS but not by lyso-PE (Fig. 7B). This result indicates that PS is essential for Cryptococcus viability.

**Cho1 is essential for normal mitochondrial function**

Mitochondria are essential organelles as they supply the cell with energy. Furthermore, mitochondria protect the cell against reactive oxygen species (ROS), as disruption of the electron transport chain leads to an increase in ROS, which in turn damages nucleic acids, proteins, and lipids, eventually leading to cell death (28, 29). Because the mitochondrial membrane is composed of lipids, including PS, PE, and PC, and Sc Cho1 has been shown to play a role in mitochondria function (30, 31), we hypothesized that **Cn Cho1** may play an important role in mitochondrial function by contributing to lipid homeostasis in the mitochondrial membrane. To test our hypothesis, we first used MitoTracker Red FM to examine whether reduced levels of **Cn Cho1** expression perturb mitochondrial membrane potential. The results by confocal microscopy showed that in the presence of CuSO₄, **PCTR4-CHO1** strain was unable to retain the MitoTracker Red FM, whereas in the presence of BCS or SC medium alone, this dye accumulated in the mitochondria (Fig. 8A and Fig. S4). To determine whether **PCTR4-CHO1** strain lacked mitochondria, yeast mitochondrial DNA (mtDNA) was examined using DAPI. Mitochondrial DNA staining using DAPI indicated that this mitochondrial dysfunction was not due to the lack of mitochondria or damage to mtDNA, as **PCTR4-CHO1** strain treated with CuSO₄ still showed the mtDNA signal (Fig. 8B and Fig. S4). We also determined the cell viability after CuSO₄ treatment for 0, 2, and 4 h by measuring the yeast CFU of treated cells, and we found that the majority of the cell populations were still viable (Table S2). These data support our hypothesis that Cho1 is essential to maintain healthy mitochondria by helping maintain mitochondrial membrane integrity.

Perturbed mitochondrial respiratory chain may lead to increased generation of ROS (29). Therefore, we used the ROS indicator dye H₂DCFDA to monitor ROS generation. After a 2-h incubation in the presence of CuSO₄, BCS, or SC medium alone, there was no significant change in ROS generation in **PCTR4-CHO1** and H99 strain. However, the ROS level significantly increased in the **PCTR4-CHO1** strain as compared with H99 after 4 and 6 h of incubation (Fig. 8C). These data demonstrate that **Cn Cho1** plays an important role in **C. neoformans** mitochondrial function, and mitochondrial dysfunction likely contributes to the lethality of **Cn Cho1**.

**Discussion**

In this study, we have identified and characterized the **Cn Cho1** protein, the sole homolog of PS synthase homolog in **C. neoformans**. We have shown that heterologous expression of **Cn Cho1** in the **S. cerevisiae cho1Δ** mutant fully complements the PS synthase activity, confirming that **Cn Cho1** is a PS synthase. We further found that the **Cryptococcus** PS synthase is essential for fungal viability, despite the existence of an active Kennedy pathway to produce PE and PC. Our results demonstrated that PS is essential for fungal cell viability, and although the main reason for **Cn Cho1** essentiality is not fully understood, we did show that it contributes to mitochondrial function.

Fungal Cho1 homologs have been shown to play an important role in cell signaling and synthesis of major cellular phospholipids, but this is the first report indicating that Cho1 is essential for cell viability in a fungus. In **S. cerevisiae**, the **cho1Δ** mutant is viable but auxotrophic for ethanolamine. The **S. cerevisiae** **cho1Δ** mutant grows normally in the presence of ethanolamine or choline, suggesting that the Kennedy pathway can support lipid production in this organism (32). The **cho1Δ/cho1Δ** null mutant is also viable in **C. albicans** (15). Interestingly, although the in vitro growth of this mutant can be rescued by addition of ethanolamine, Cho1 is essential for **Candida** virulence in a murine infection model and has thus been proposed as a potential drug target (15, 33). A recent study further demonstrated that the mutant cells might not be able to acquire sufficient ethanolamine from the host, leading to the avirulent phenotype (34). Why Cho1 is essential in **C. neoformans** remains unclear. Our 1D-TLC analysis showed that PE and PC can be produced by cells supplemented with choline or ethanolamine in culture medium, indicating that the Kennedy pathway remains active in **C. neoformans**. These data suggest that the essentiality of Cho1 in **C. neoformans** is not due to a lack of PE and PC production via the Kennedy pathway, although we do not know how much of these lipids are synthesized through this pathway in this fungus.

Using the **CTR4**-inducible promoter system, we were able to knock down the PS synthase activity to a level that prevents cell growth. The growth arrest of the **PCTR4-CHO1**-expressing strain in the presence of CuSO₄ was rescued by the addition of lyso-PS, but not lyso-PE, indicating that PS is essential for **C. neoformans** and that the inability to produce PS is the reason for **Cn cho1Δ** mutant lethality. This is consistent with the conclusion that **C. neoformans** can make PE and PC via the Kennedy pathway but that PS has to be synthesized using PS synthase Cho1. PS can be converted to PE by PS decarboxylase Psd1 and Psd2. This step is generally considered irreversible in fungi such as **S. cerevisiae** and is likely to be irreversible in **C. neoformans** as well.

PS plays several important cellular roles. Besides being a key component of the lipid bilayer membrane, PS also functions as a phagocytosis-inducing signal for macrophages and a signal to trigger inflammatory activity during infection (8, 10, 11). Because the **Cn cho1Δ** mutant is inviable, we could not test the function of Cho1 in host–pathogen interactions. Based on our results, it is likely that PS is an essential part of the cell membrane in **C. neoformans**. The importance of PS in **C. neoformans** is also underscored by the phenotype of the lipid flippase **cdc50A** mutant, which shows PS accumulation on the extracytoplasmic leaflet of the membrane. This mutant is also defective for fungal virulence and shows increased phagocytosis and killing by macrophages in vitro (19).

Another potential role of Cho1 is to maintain mitochondrial integrity and function. The inner and outer yeast mitochondrial membranes are composed of various lipids, including PC, PE, PS, phosphatidylinositol, phosphatidic acid, phosphatidylglyc-

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erol, and cardiolipin. Two abundant phospholipids, PC and PE, are mainly synthesized from PS (29–31). Although distributions of these lipids in the inner and outer mitochondrial membranes differ, a change in mitochondrial lipid homeostasis can lead to mitochondrial respiration defects (31). In S. cerevisiae, the loss of PS or PE biosynthesis leads to the formation of petite mutants, which cannot utilize nonfermentable carbon sources such as glycerol due to mitochondrial defects (29, 31). In contrast, C. albicans is a nonpetite forming yeast, indicating that it cannot grow without functional mitochondria (15). In C. neoformans, mutants directly involved in mitochondrial function display disturbed mitochondrial action potential and increased levels of intracellular ROS (28). Functional mitochondria are required for the survival of C. neoformans, especially under low-oxygen conditions. Our results indicate that Cho1 plays an important role in maintaining the integrity of the mitochondrial...
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Table 1
Strains used in this study

| Strains used in this study | Genotype | Source/Ref. |
|---------------------------|----------|------------|
| C. neoformans strains     |          |            |
| H99                       | MATα wildtype | 43         |
| AI187                     | MATα/a diploid wildtype | 24 |
| CUX663                    | A1187 cho1Δ:NEO CHO1 | This study |
| CUX664                    | MATα P<sup>C</sup>C7-CHO1mCherry-NAT | This study |
| CUX1081                   | MATα cho1Δ:NEO P<sup>C</sup>CTR4-CHO1-NAT | This study |
| CUX1082                   | MATα cho1Δ:NAT | UCSF 2015 deletion collection |
| S. cerevisiae strains     |          |            |
| BY4741                    | MATα his3Δ1 leu2Δ0 ura3Δ0 | ATCC deletion collection |
| W303-1A                   | MATα ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 | 44 |
| JSY94A                    | MATα ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 cho1Δ:TRP1 | 41 |
| YUX104                    | MATα ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 | This study |
| YUX105                    | MATα ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 cho1Δ:TRP1 | This study |
| YUX106                    | MATα ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 | This study |

Heterologous expression of Cn Cho1 into Sc cho1Δ mutant

C. neoformans Cn CHO1 cDNA (CNAG_01469) was cloned into yeast expression vector pTH19 (P<sup>ADH1</sup> URA3 2μ) (35) and transformed in S. cerevisiae cho1 mutant (Sc cho1Δ) background, in which the expression of the Cn CHO1 gene was under the control of the ADH1 promoter. Cultures of S. cerevisiae strains WT W303-1A, Sc cho1Δ expressing empty vector, and Sc cho1Δ-expressing Cn CHO1 gene (Sc cho1Δ + Cn CHO1) (Table 1) were prepared with serial dilutions on SC-Ura medium with and without 1 mM ethanolamine. The results were photographed after 48 h of incubation at 30 or 37 °C.

Generation of the cho1Δ mutant strain

Generation of a CHO1 deletion mutant in haploid WT H99 and diploid strain AI187 backgrounds was attempted following the strategy illustrated in Fig. 5A. The cho1Δ mutant was generated by overlap PCR and biolistic transformation as described previously (36). The 5'- and 3'-flanking regions of the CHO1 gene were amplified from H99 genomic DNA with primers CX862/CX863 and CX864/CX865, respectively (Table S1). The dominant selectable marker (NEO) was amplified with the M13 primers (M13F and M13R) from plasmid pAF1 (37). Each target gene replacement cassette was generated by overlap PCR with primers CX862/CX865 (Table S1). Purified overlap PCR products were precipitated onto 10-μl gold microcarrier beads (0.6 μm; Bio-Rad), and biolistically transformed in H99 and AI187 as described previously (38). Stable transformants were selected on YPD medium containing 448 (200 mg/liter). To screen for mutants of the CHO1 gene, diagnostic PCR was performed by analyzing the removal of Cn CHO1 ORF with primers CX866/CX867 (Table S1). Positive transformants identified by the PCR screen were further confirmed with primers CX868/JH8994.

Generation of the P<sup>CTR4</sup>-CHO1-expressing strain

The native CHO1 promoter was replaced by the CTR4-inducible promoter following the strategy illustrated in Fig. S2. The fragment containing CHO1 ORF and the downstream terminal region was amplified from H99 genomic DNA with primers CX1269/CX1270 (Table S1). Amplified PCR product was cloned into the BamHI sites of the pCTR4 vector (kindly provided by Dr. Tamara Doering laboratory, Washington University in St. Louis (25)) using an In-Fusion HD cloning kit (Takara, CA). The CHO1 upstream and selectable marker (NAT) fragments were amplified using primers CX1271/CX1272 and CX5/CX1273, respectively (Table S1). Overlap PCR product (CHO1 upstream-NAT fragment) was cloned into Ndel sites of CHO1 ORF pCTR4-2 vector by In-Fusion HD cloning. The constructed vector was biolistically transformed into WT H99. Positive transformants identified by PCR using CX1306/JH8994 and CX1307/CX1276 primers (Table S1) were further confirmed by Southern blotting and gene sequencing.
**RNA isolation and cDNA synthesis**

H99 and H99-expressing P<sub>CTR4</sub>-CHO1 strains were grown in 50 ml of YPD medium at 30 °C overnight. Cells were then centrifuged, washed twice with sterile water, and transferred into 50 ml of SC medium containing either 200 μM BCS for inducing condition or 25 μM or 1 mM CuSO₄ with 1 mM ascorbic acid for suppressing condition. Cell cultures were incubated at 30 °C for 20 h. RNAs were isolated using the TRIzol method followed by phenol/chloroform extraction. The genomic DNA was eliminated using rDNase and RNA purification kit (Takara, CA). The quality of RNA was confirmed by measuring the A<sub>260/280</sub> m ratio on a spectrophotometer (Bio-Rad) and visualizing on an RNA gel. For cDNA synthesis, RNA samples were reverse-transcribed using SMARTScribe reverse transcriptase (Takara, CA).

To determine CHO1 gene expression in H99 under various inositol concentrations, cells were grown in 50 ml of YPD medium at 30 °C overnight. Cells were then centrifuged, washed twice with sterile water, and transferred into 50 ml of SC supplemented with 0.1 or 1% inositol (w/v), respectively. Cells were collected at two time points, 4 h after incubation at 30 °C, and RNA was isolated as described above.

**Real-time PCR**

The real-time PCR was performed using 2× master mix SYBR Green on an Mx4000 quantitative PCR system (Stratagene, CA). The amplification conditions consist of 1 cycle at 95 °C for 10 min, 40 cycles at 95 °C for 30 s, 55 °C for 30 s, and 72 °C for 30 s, and 1 cycle at 95 °C for 1 min, 55 °C for 30 s, and 95 °C for 30 s. The results were normalized to GAPDH and analyzed using 2<sup>-ΔΔCt</sup> method (39).

**Protein extraction and Western blot analysis**

A previously described method with minor modification was used to determine whether CHO1 is an integral membrane protein (40). C. neoformans strain expressing CHO1:mCherry (Table 1) was grown in 50 ml of YPD to early mid-log phase before harvesting by centrifugation at 1500 × g for 5 min at 4 °C. Cells were washed with cold water and centrifuged at 1500 × g for 5 min. Cells were then resuspended in 20 ml of Tris-HCl/EDTA buffer (0.1 M Tris-HCl, pH 7.5, 0.5 M EDTA) with 1 mM DTT and incubated at 30 °C for 2 h. Cell pellets were washed with 1 M sorbitol and incubated in 1 M sorbitol with lyticase (1 μg/μl) and chitinase (30 mg/ml) at 30 °C for 3 h. One gram of spheroplasts was resuspended in homogenizing buffer (150 mM NaCl, 20 mM Tris-HCl, pH 8) and lysed using a beads beater (8 times for 40 s each cycle). Cells were centrifuged at 13,000 × g for 10 min at 4 °C. The insoluble pellet was resuspended in 8 ml of either buffer alone (150 mM NaCl, 20 mM Tris, pH 8.0, 5 mM MgCl₂, 1 mM phenylmethylsulfonyl fluoride), buffer with 1% Triton X-100, or buffer with 1 M NaCl. Extracts were incubated on ice for 30 min and then separated into a soluble and pellet fraction by ultracentrifugation at 226,000 × g for 60 min at 4 °C. Equivalent amounts from the pellet and the supernatant were analyzed by Western blotting and SDS-PAGE. The antibodies were used at the following dilutions: RFP-tag antibody 1:1000 (GenScript) and anti-rabbit 1:1000 (GenScript).

**Lipid extraction, separation, and identification**

Lipid isolation followed a previously reported method with modifications (41, 42). S. cerevisiae WT strain BY4742, C. neoformans WT strain H99, and its mutant of the EKI/CKI kinase homolog (Cn cki1Δ) strains (Table 1) were grown in YPD overnight to A<sub>600 nm</sub> = 5. Cells were centrifuged, washed twice in sterile water, and incubated at a starting A<sub>600 nm</sub> = 0.1 in 2 ml of SC medium with 10 μCi of [14C]ethanolamine and 10 μCi of [3H]choline overnight at 30 °C. Labeled cells were harvested by centrifugation, washed twice with sterile water, and incubated in 1 ml of spheroplast buffer (1 M sorbitol, 0.05 M sodium phosphate monobasic, 0.1% 2-mercaptoethanol (v/v)) with 100 μg/ml chitinase for 1 h at 30 °C. Cells were collected by centrifugation, washed with sterile water, and resuspended in chloroform/methanol (2:1). Glass beads were added, and cells were lysed using a beads beater (8 times for 40 s each cycle). Lysate was transferred into new 1.5-ml tubes and incubated on a shaker at room temperature for 30 min. To separate the organic phase containing phospholipids from the aqueous phase containing nonlipid cellular material, 0.2 volume of water was added. Mixture was vortexed and centrifuged at 1000 × g for 1 min. Eighteen microliters of lipids were loaded onto HPTLC silica gel (Millipore, MA). Solvent used in the mobile phase consisted of chloroform/methanol/acetic acid (65:25:10). Lipids were visualized using 99.99% iodide and autoradiography film (Denville Scientific, NJ).

**ER tracker staining**

C. neoformans overnight cultures were washed twice in Hanks’ balanced salt solution (HBSS) buffer (1.26 mM CaCl₂, 5.33 mM KCl, 0.44 mM KH₂PO₄, 0.5 mM MgCl₂·6H₂O, 0.44 mM MgSO₄·7H₂O, 138 mM NaCl, 4 mM NaHCO₃, 0.3 mM Na₂HPO₄, and 5.6 mM D-glucose) before being fixed with 3.7% formaldehyde for 5 min at 30 °C. Fixed cells were washed with HBSS buffer three times and stained with 1 μM ER tracker (Life Technologies, Inc.) for 30 min at 30 °C before being observed under the fluorescence microscope (Nikon).

**Mitochondria and DNA staining**

C. neoformans overnight cultures were resuspended in SC medium containing either 200 μM BCS or 1 mM CuSO₄ and 1 mM ascorbic acid. Cells were incubated at 30 °C for 6 h and then washed twice in 1× PBS and resuspended in 1× PBS with 0.2 μM MitoTracker Red FM (Invitrogen) and 1 μg of DAPI (ThermoFisher Scientific). Mixture was incubated for 10 min at 30 °C. Cells were then washed twice with 1× PBS and observed under a fluorescence microscope (Nikon).

**ROS measurement**

ROS detection technique followed a previously reported method (28). Cells were grown overnight in YPD medium at 30 °C. The following day, cells were diluted in SC medium and allowed to grow until A<sub>600 nm</sub> reached 0.5. H₂DCFDA (Invitrogen) at a final concentration of 10 μM was added, and cells were incubated for an additional 2 h. Cells were then washed to remove the excess dye and resuspended in SC, SC with 200 μM BCS, and SC with 1 mM CuSO₄ to further incubate for 2, 4, and
Phosphatidylserine synthesis is essential in Cryptococcus

6 h. Cells were then harvested, washed with 1 × PBS, and resuspended in 1 ml of PBS. Fluorescence signal was analyzed using Accuri Flow Cytometer (BD Biosciences). Two-tail t test was used for statistical analysis, where * represents \( p < 0.05 \), and ** represents \( p < 0.01 \).

PS synthase activity assay

All assays were conducted at 30 °C in a total volume of 0.1 ml. PS synthase activity was measured following the incorporation of water-soluble \([3-3H]\)serine (10,000 cpm/nmol) into chloroform-soluble \([3-3H]PS\) (16, 22, 23). The enzyme reaction contained 50 mM Tris-HCl, pH 8.0, 0.6 mM MnCl₂, 4 mM Triton X-100, 0.2 mM CDP-DAG, and 0.5 mM serine (41). The enzyme assays were conducted in triplicate, and the average S.D. of the assays was ±5%. The reactions were linear with time and protein concentration. A unit of PS synthase activity was defined as the amount of enzyme that catalyzed the formation of 1 nmol of product/min.

Statistical analysis

Means with standard deviations were calculated using the t test for multiple comparisons. The value of \( p < 0.05 \) was considered a significant difference.

Author contributions—P. K., Y. W., G.-S. H., K. J. G., and C. X. data curation; P. K., Y. W., G.-S. H., K. J. G., Y.-G. G., and C. X. formal analysis; P. K., Y. W., and C. X. validation; P. K., Y. W., G.-S. H., K. J. G., Y.-G. G., and C. X. investigation; P. K., Y. W., and C. X. visualization; P. K., Y. W., G.-S. H., K. J. G., and C. X. methodology; P. K. and C. X. writing-original draft; P. K., Y.-G. G., M. C., and C. X. project administration; P. K., G.-S. H., Y.-G. G., M. C., and C. X. writing-review and editing; K. J. G., Y.-G. G., and C. X. software; G. M. C. and C. X. conceptualization; G. M. C. and C. X. supervision; G. M. C. and C. X. funding acquisition.

Acknowledgments—We thank Erika Shor for critical reading and editing of the manuscript and valuable comments for the study. We thank Dr. Xilin Zhao for valuable comments and material support for the ROS experiments. We thank Dr. Todd Reynolds for valuable comments and material support for the C. neoformans genome. We also acknowledge use of the C. neoformans genome project-soluble \([3-3H]PS\) (16, 22, 23).

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