Multiple Alternative Carbon Pathways Combine To Promote *Candida albicans* Stress Resistance, Immune Interactions, and Virulence

Robert B. Williams,* Michael C. Lorenz*

*Department of Microbiology and Molecular Genetics, University of Texas McGovern Medical School and the MD Anderson Cancer Center UTHealth Graduate School of Biomedical Sciences, Houston, Texas, USA

**ABSTRACT** The phagocytic cells of the innate immune system are an essential first line of antimicrobial defense, and yet *Candida albicans*, one of the most problematic fungal pathogens, is capable of resisting the stresses imposed by the macrophage phagosome, eventually resulting in the destruction of the phagocyte. *C. albicans* rapidly adapts to the phagosome by upregulating multiple alternative carbon utilization pathways, particularly those for amino acids, carboxylic acids, and *N*-acetylglucosamine (GlcNAc). Here, we report that *C. albicans* recognizes these carbon sources both as crucial nutrients and as independent signals in its environment. Even in the presence of glucose, each carbon source promotes increased resistance to a unique profile of stressors; lactate promotes increased resistance to osmotic and cell wall stresses, amino acids increased resistance to oxidative and nitrosative stresses, and GlcNAc increased resistance to oxidative stress and caspofungin, while all three alternative carbon sources have been shown to induce resistance to fluconazole. Moreover, we show mutants incapable of utilizing these carbon sources, in particular, strains engineered to be defective in all three pathways, are significantly attenuated in both macrophage and mouse models, with additive effects observed as multiple carbon pathways are eliminated, suggesting that *C. albicans* simultaneously utilizes multiple carbon sources within the macrophage phagosome and during disseminated candidiasis. Taking the data together, we propose that, in addition to providing energy to the pathogen within host environments, alternative carbon sources serve as niche-specific priming signals that allow *C. albicans* to recognize microenvironments within the host and to prepare for stresses associated with that niche, thus promoting host adaptation and virulence.

**IMPORTANCE** *Candida albicans* is a fungal pathogen and a significant cause of morbidity and mortality, particularly in people with defects, sometimes minor ones, in innate immunity. The phagocytes of the innate immune system, particularly macrophages and neutrophils, generally restrict this organism to its normal commensal niches, but *C. albicans* shows a robust and multifaceted response to these cell types. Inside macrophages, a key component of this response is the activation of multiple pathways for the utilization of alternative carbon sources, particularly amino acids, carboxylic acids, and *N*-acetylglucosamine. These carbon sources are key sources of energy and biomass but also independently promote stress resistance, induce cell wall alterations, and affect *C. albicans* interactions with macrophages. Engineered strains incapable of utilizing these alternative carbon pathways are attenuated in infection models. These data suggest that *C. albicans* recognizes nutrient composition as an indicator of specific host environments and tailors its responses accordingly.

**KEYWORDS** *Candida*, macrophage, virulence, carboxylic acid, amino acid, *N*-acetylglucosamine, metabolism, stress, fungal pathogenesis, nutritional immunity, host-pathogen interactions
Candida albicans is both a common human commensal and a serious clinical problem, capable of causing an array of mucosal and invasive infections. The most severe form of infection, disseminated candidiasis, is the most common fungal nosocomial bloodstream infection (1, 2). C. albicans often infects the most vulnerable patient populations; those undergoing chemotherapy or organ transplantations or with implanted medical devices are especially vulnerable to disseminated candidiasis due to prolonged hospital stays and weakened immune systems (3). With mortality rates reaching 40% and an increasing fear of antifungal resistance, it is critical that we understand the basis of the many pathogenesis mechanisms that C. albicans utilizes during disseminated infection.

The phagocytes of the innate immune system are among the first defense mechanisms deployed to combat a Candida infection (4). Neutrophils and macrophages typically clear pathogens by bombarding the organism with a variety of stressors such as oxidative stress, nitrosative stress, and hydrolytic enzymes, some of which are potentiated by phagosome acidification (5–9). C. albicans, however, is capable of surviving macrophage phagocytosis by resisting these stressors, actively neutralizing the pH of the phagosome, and ultimately shifting its morphology to the hyphal form. These stresses lead to the death of the macrophage via pyroptosis and allow the fungal cell to escape the phagocyte and continue dissemination (10–17). Thus, we and others have characterized the responses that allow C. albicans to adapt and overcome the phagosomal environment. Transcriptomic studies reveal that C. albicans rapidly adapts to the macrophage phagosome by initially shifting its metabolism away from glycolysis in favor of alternative carbon sources such as carboxylic acids, amino acids, peptides, N-acetylglucosamine (GlcNAc), and fatty acids (10, 18–24). These compounds are sources of energy and biomass, and mutations that impair utilization of these nutrients are attenuated in macrophage and mouse virulence models.

Beyond their role as nutrients, three of these carbon sources also directly contribute to C. albicans-phagocyte interactions: carboxylic acids such as lactate, amino acids, and GlcNAc. Catabolism of these alternative carbon sources results in a significant increase in extracellular pH, both in vitro and within the phagososome, but the physiological effects diverge from there (10, 13, 14). The presence of lactate increases C. albicans resistance to a variety of host-relevant stressors, enhances biofilm formation, and reduces recognition by macrophages (15, 25–28). These adaptations are mediated in part by alterations in the thickness and composition of the cell wall (15, 26, 27). Amino acids and GlcNAc strongly induce hyphal morphogenesis, contributing to fungal survival both within macrophages and during disseminated candidiasis (10–12). How the presence of amino acids or GlcNAc might affect stress resistance, cell wall morphology, or phagocyte recognition has not previously been addressed.

The genetic requirements for the physiological changes promoted by the different carbon sources are also distinct, which has allowed the dissection of the roles of the individual alternative carbon pathways (10, 13, 14, 26, 29, 30). JEN1 and JEN2 encode the carboxylic acid transporters (particularly mono- and dicarboxylic acids such as lactate, pyruvate, α-ketoglutarate [α-KG], and malate) that are strongly induced after phagocytosis by macrophages or neutrophils at the protein and RNA level, but the double deletion jen1Δ jen2Δ strain, which is incapable of importing several carboxylic acids, has been reported to show no defects in a disseminated candidiasis mouse model (18, 31, 32). Mutants lacking STP2, encoding a transcription factor that regulates amino acid uptake and utilization, are unable to neutralize the acidic environment and are modestly attenuated in both macrophage and mouse models (10). Mutants incapable of catabolizing GlcNAc are likewise modestly attenuated in macrophages and mice (13, 33).

The modest individual phenotypes led us to hypothesize that C. albicans simultaneously utilizes all of these nutrients during infection, with each carbon source providing unique benefits to the pathogen. Combining these effects through genetic elimination of multiple carbon pathways would thus have additive effects on fitness in macrophage interactions and disseminated infections. In support of this hypothesis, we
report here that alternative carbon sources serve as unique priming signals with respect to the host environments that C. albicans may encounter. Each alternative carbon source induces a unique pattern of protection from host-relevant stressors compared to glucose-grown cells. Exposure to these nutrients also alters interactions with immune cells, likely as a result of significant changes in the fungal cell wall. Additionally, mutant strains lacking multiple carbon utilization pathways are significantly attenuated in macrophages and in mouse models. We conclude that C. albicans makes use of multiple alternative carbon sources simultaneously and that these serve as more than energy sources for the pathogen, each conferring unique benefits that allow it to adapt and cause infection in a variety of host environments.

RESULTS

Alternative carbon sources affect C. albicans stress tolerance. Abundant evidence suggests that C. albicans utilizes multiple carbon sources in vivo, including glucose, carboxylic acids (lactate, α-ketoglutarate), amino acids, and GlcNAc, and that the catabolism of these compounds has effects beyond generation of energy and biomass (10, 13, 14, 18, 22, 26–28, 33). In particular, lactate induces changes to C. albicans cell wall morphology, stress resistance, and immune interactions that promote pathogenesis (15, 25–27). We asked whether these physiological adaptations might be triggered by additional alternative carbon sources, and whether the responses were carbon source specific.

C. albicans encounters multiple stressors, including oxidative, nitrosative, osmotic, cell wall, and antifungal stresses, during disseminated infection. To assess how alternative carbon utilization affects host-relevant stress resistance, C. albicans cells were pregrown in minimal yeast nitrogen base with allantoin (YNBA) media containing 1% glucose or containing 1% glucose plus 1% of one of the three alternative sources, namely, lactate, Casamino Acids (CAA), or GlcNAc. Cells were then transferred to fresh media with or without the indicated stress. Growth was assessed in multiwell plates over time at 37°C in a multifunctional plate reader and quantified by assessing the change in optical density at 600 nm (OD600), relative to the no-stress control, after 8 h (Fig. 1; representative growth curves are presented in Fig. S1 in the supplemental material). In agreement with previous reports (15, 27), lactate protected C. albicans from high concentrations of the cell wall stressors calcofluor white (CFW) and Congo red (Fig. 1A and B). In contrast, lactate did not alter sensitivity to oxidative or nitrosative stresses (Fig. 1C to E). Lactate induced protection from the antifungal drug fluconazole but not from caspofungin (Fig. 1F and G). Similarly to previous reports, lactate also increased resistance to salt stress (Fig. 1H) (15).

Interestingly, amino acids and GlcNAc also induced resistance to some of these stresses, but in distinct patterns: amino acids strongly induced protection from oxidative stressors tert-butyl-hydroperoxide (TBO) and H2O2 (Fig. 1C and D) and more modestly induced protection from nitrosative stress and CFW (Fig. 1A and H). It had no effect on Congo Red or nitrosative or osmotic stress (Fig. 1B, E, and H). In contrast, GlcNAc enhanced resistance to TBO but not peroxide (Fig. 1C and D) and had no effect on nitrosative, cell wall, or osmotic stress (Fig. 1A, B, E, and H). All three alternative carbon sources promoted fluconazole resistance, whereas amino acids and GlcNAc, but not lactate, protected cells from caspofungin (Fig. 1F and G).

As a corollary, we also asked whether alternative carbon sources provided protection against acute toxicity after shorter-term exposure to extreme stress concentrations. The proportion of cells grown under all three conditions that survived exposure to 10 mM TBO was much greater than that seen with glucose alone (Fig. S2). In contrast, only lactate protected against 2 M NaCl. Together, these data suggest that the changes in stress resistance represented an active response to the presence of these compounds and not a consequence of glucose deprivation. Moreover, each alternative carbon pathway was found to be physiologically distinct, inducing different patterns of stress resistance, which raises the possibility that nutritional composition is a signal that may allow C. albicans to anticipate likely stresses in specific host environments.
Alternative carbon utilization promotes resistance to a diverse array of stressors. SC5314 was grown for 6 h in minimal media containing 1% glucose or containing 1% glucose plus 1% of the indicated alternative carbon source and then transferred to the same media with or without the indicated stressor in a 96-well plate and incubated at 37°C with aeration in a plate reader for 16 h. The data represent the difference in the change in OD600 at the 8-h time point, expressed as a ratio of the level seen the presence of stress to that seen in the absence of stress. N = 5 for each stress condition. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001 (unpaired t tests).
Alternative carbon sources alter interactions with macrophages. Lactate exposure alters interactions with macrophages, but little is known about the effects of amino acids and GlcNAc (27). *C. albicans* was again pregrown to mid-log phase in minimal media containing glucose with or without the alternative carbon sources before assessing macrophage interactions. Media were buffered to pH 5.5 to prevent the cell wall changes that can occur under conditions of pH fluctuations (30). As previously reported, the presence of lactate modestly decreased the rate at which cells associated with and were phagocytosed by macrophages (Fig. 2A and B). Interestingly, pregrowth in amino acids resulted in increased phagocytosis rates compared to those seen with glucose-grown cells, while GlcNAc had no significant effect. Despite the increased phagocytosis, the amino acid-grown cells showed significantly improved fungal survival compared to the lactate-exposed cells, but the differences from the glucose-grown cells were not statistically significant (Fig. 2C). Together, these data show that alternative carbon sources, particularly lactate and amino acids, distinctly affect immune recognition of fungal cells.

*C. albicans* occupies a phagolysosome of significantly higher pH than would be expected, and we have previously found that this is associated with utilization of alternative carbon sources (10, 14, 34, 35). We next asked whether prior exposure to these nutrients would affect the ability to neutralize the pH of macrophage phagosomes. To address this issue, we loaded macrophages with the acidophilic dye Lyso-Tracker red prior to culturing with *C. albicans* cells (pregrown as described above) in media containing glucose, lactate, amino acids, or GlcNAc. The level of phagosome acidity immediately surrounding the fungal cell was calculated by measuring Lyso-Tracker intensity immediately outside the fungal cell, as previously described (11, 13). Interestingly, no differences were observed in the levels of phagosome acidity after exposure to alternative carbon sources (Fig. S3).

Alternative carbon sources alter cell wall structure. The changes observed in response to the presence of lactate were found to be associated with alterations to cell wall structure (15). To assess whether amino acids and GlcNAc had similar effects on the cell wall, we visualized this organelle using transmission electron microscopy (Fig. 3). *C. albicans* was grown to mid-log phase in minimal medium (YNBA) with a carbon source (glucose, Casamino Acids, lactate, or GlcNAc) (1%) buffered to pH 5.5 before fixing and processing. The diameter of yeast-phase cells grown on alternative carbon sources was significantly reduced compared to that of glucose-grown cells (Fig. 3A). This is in line with observations in both single-celled and multicellular organisms indicating that nutrient restriction generally results in reduced cell size (36–39). Growth in alternative carbon sources reduced the overall thickness of the cell wall (Fig. 3B); the inner glucan layer was markedly thinner, while the outer mannann layer was mostly unchanged (Fig. 3C). As a result, the ratio of the thicknesses of the glucan and mannann layers was greatly reduced in cells grown in the absence of glucose (Fig. 3D). Notably, the glucan layer of both amino acid-grown and GlcNAc-grown cells was more electron dense than that of glucose-grown and lactate-grown cells (Fig. 3E). Electron-dense granules of 30 to 100 nm were frequently seen in the cell walls of glucose-grown cells but not under the other conditions (Fig. 3E); we assume that they represent extracellular vesicles transiting the wall. Thus, utilization of lactate, GlcNAc, and amino acids results in distinct decreases in cell size and cell wall thickness, shifts in the ratios of the thicknesses of the glucan and mannann layers, and alterations in cell wall density.

Strains unable to utilize alternative carbon sources are defective in multiple fitness assays. Since exposure of *C. albicans* to individual alternative carbon sources has overlapping effects on cell wall morphology, stress resistance, and interactions with macrophages, we next asked whether genetically interfering with alternative carbon utilization would decrease fitness within the host. We conducted a genetic screen for mutants that grow poorly on media containing lactate as the sole carbon source, using available libraries of ~850 strains (the Noble and Homann libraries) (40, 41). The libraries were grown in yeast extract-peptone-dextrose (YPD) medium and transferred...
FIG 2 Alternative nutrients affect *C. albicans* interactions with macrophages. SC5314 was grown for 6 h in minimal media containing 1% of the indicated carbon source prior to each experiment. (A) SC5314 expressing scarlet fluorescent protein was coincubated with J774A macrophages stained with nucBlue and actin green in DMEM at 5% CO₂ and 37°C and was imaged every 15 min in an imaging plate reader. Cell association was quantified as the number of red *C. albicans* cells that were overlaid with green macrophages. n = 5. (B) Similarly to the method described for panel A, scarlet fluorescent protein-expressing SC5314 was coincubated with J774 macrophages stained with DAPI (4′,6-diamidino-2-phenylindole) for 60 min. Samples were fixed with paraformaldehyde and counterstained with FITC to identify external fungal cells. n = 5. (C) Fungal survival was assessed after 16 h of coincubation with J774 macrophages using a modified CFU assay. Percent survival data are compared to data from a control well containing *C. albicans* strains incubated without macrophages. n = 5. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001 (unpaired t tests).
to minimal medium containing 1% lactate (plus 10 mM arginine, as the library strains are arginine auxotrophs) and assessed for growth and pH neutralization using the pH indicator bromocresol purple. Hits from this screen were combined with those from a previously reported screen for mutants impaired on the same media with α-ketoglutarate (10). Mutants with initial growth defects were confirmed via PCR and converted to prototrophy using Cip30, and doubling times were assessed in multiple media (Fig. 4).

These mutants had a spectrum of phenotypes (Fig. 4), with some strains impaired in all carbon sources and others in a subset. A few strains (ali1Δ, cox4Δ, sin3Δ mutants) also conferred significant growth defects in glucose-containing minimal media, as has been reported previously (40, 42). Unsurprisingly, many of the mutants encoded proteins associated with functions required for aerobic respiration. These include Cox4 (a subunit of cytochrome c), Ali1 (a membrane-bound NADH-ubiquinone oxidoreduc-
tase), and Oar1 (a putative mitochondrial protein whose homologs have NADPH-dependent 3-oxoacyl reductase activity) (40, 42). Two mutants have known roles in other aspects of carbon metabolism, including the Snf1 complex protein Kis1 and Pex13, which is required for protein import to the peroxisome (43–45). Others had no clear connection to carbon utilization, including Pep8 (a protein involved in endosome-to-Golgi transport), Swi4 (part of the SBF complex required for the G1/S transition), and Sin3 (a global transcriptional corepressor) (46–49). The well-characterized morphology regulator Cph1 also has defects in growth on alternative carbon sources, though this particular library mutant also grew more slowly when glucose was present, which is not a phenotype that was previously associated with the *cph1*Δ mutation. Notably, the *cox4*Δ/H9004 mutant was able to grow in GlcNAc but did so without forming hyphae.

These mutants were evaluated for fitness defects in multiple macrophage assays (Fig. 5). Since alternative carbon sources promote extracellular pH neutralization, we assessed the ability of the mutants to neutralize the phagosome. Several mutants occupied significantly more acidic phagosomes within J774A.1 cells, indicative of pH neutralization defects (Fig. 5A). Failure to neutralize the phagosome was previously shown to be associated with impaired fungal survival in macrophages, and, indeed, all of the screened mutants, with the possible exception of the *kis1*Δ strain (where the decrease was not statistically significant), survived a 16-h coincubation less readily than the control strain (Fig. 5B). Among these mutants, the *ali1*Δ, *cox4*Δ, and *oar1*Δ strains also showed notable hyphal defects within the macrophage. The *cox4*Δ mutant appeared to be entirely incapable of forming hyphae, as it was found in the yeast and pseudohyphal forms under conditions that strongly induced hypha (RPMI plus serum, 5% CO₂, 37°C) (data not shown). We also tested macrophage survival, using release of cytosolic lactate dehydrogenase (LDH) as a proxy for membrane damage, and found that all of the mutants, with the possible exception of the *pex13*Δ and *swi4*Δ strains, caused less damage to macrophages (Fig. 5C). Thus, among multiple mutants with defects in utilization of alternative carbon sources, nearly all were attenuated in multiple aspects of interactions with macrophage, further indicating the importance of alternative carbon utilization during infection.

**Engineered carbon-deficient mutants are impaired in multiple virulence models.** As a second and more direct approach to address this issue, we employed the CRISPR-Cas9 system to generate strains lacking one or more of the genes that have been previously shown to directly contribute to nutrient import and catabolism, i.e., *STP2, JEN1, JEN2, HXX1, NAG1*, and *DAC1* (11, 29, 32). Stp2 is a transcription factor that controls amino acid import and catabolism. Jen1 and Jen2 are mono- and dicarboxylic acid transporters, respectively. Hxx1, Nag1, and Dac1 are three enzymes essential for GlcNAc catabolism and are encoded by a cluster of adjacent genes on chromosome 6 such that they could be knocked out in tandem (“h-dΔ”) (50). Individually, these
mutants were previously shown to have modest impacts on fungal survival in macrophages and virulence in a disseminated mouse model (13, 14, 32). We hypothesized that these phenotypes would be additive, such that elimination of multiple carbon pathways would result in markedly increased attenuation. We thus constructed the \textit{stp2Δ}, \textit{jen1Δ jen2Δ}, and \textit{h-dΔ} mutant strains and all possible combinations thereof, including the quadruple mutant, termed the “ΔΔΔΔ” strain. This strain was unable to

FIG 5 Multiple genetic screen mutants are defective within macrophages. (A) Phagosomal pH was estimated using LysoTracker Red staining as previously described (Vylkova and Lorenz (14)). In short, macrophages preloaded with acidophilic dye were coincubated with \textit{C. albicans} for 1 h and fixed, and fluorescence microscopy was performed. Red fluorescence intensity is reflective of an acidic phagosome. \textit{n} = 3. (B) Fungal survival quantified via modified CFU assay as described for Fig. 3. \textit{n} = 4. (C) Macrophage cytotoxicity was quantified via lactate dehydrogenase release after 16 h of coincubation with the indicated \textit{C. albicans} strain. Killing data are compared to results determined for chemically lysed macrophages. \textit{n} = 4. *, \textit{P} < 0.05; **, \textit{P} < 0.01; ***, \textit{P} < 0.001; ****, \textit{P} < 0.0001 (unpaired t tests).
grow on YNBA media containing lactate (monocarboxylic acid), α-ketoglutarate (dicarboxylic acid), Casamino Acids, or GlcNAc and did not neutralize the pH of these media (Fig. S4). The quadruple mutant grew at wild-type rates in YNBA-glucose or YPD media, germinated normally under conditions that strongly induced hypha (Fig. 6A; see also Fig. S5), and displayed no apparent differences in macrophage uptake rates. Additionally, supplementing the media with glycerol partially rescued growth but not pH changes, suggesting that the neutralization is dependent on alternative carbon catabolism processes (Fig. S6). Thus, we successfully generated a strain that was incapable of

FIG 6 Alternative carbon mutants are defective upon macrophage phagocytosis. (A and B) FITC-labeled strains were cocultured with J774 macrophages in DMEM for 2 h. Cells were fixed and stained with calcofluor white, and germ tube lengths were quantified in triplicate experiments (>40 per strain per replicate). Scale bar = 10 μm. (C) A LysoTracker Red assay was performed in a manner similar to that described for Fig. 5. n = 3. *, P < 0.05; **, P < 0.01; ***, P < 0.001; ****, P < 0.0001 (unpaired t tests).
using multiple alternative carbon sources but that appeared not to have pleiotropic phenotypes.

Using strains defective in one, two, and three carbon pathways (including strain ΔΔΔΔ and all intermediate strains), we assayed fitness after phagocytosis by macrophages to determine if the multiple mutants had additive defects (Fig. 6). After 2 h of coculture, each single mutant trended toward reduced hyphal lengths, while the multiple mutants showed significantly reduced hyphal elongation (Fig. 6A and B). Each carbon pathway has been associated with a reduced ability to neutralize the phagosome, and strains impaired in two or all three pathways are found in even more acidic phagosomes (Fig. 6C). Hyphal length and phagosomal pH are correlated, though it is not clear whether neutral pH induces germination or whether hyphal growth neutralizes the phagosome through membrane disruption (14, 51). These data suggest that C. albicans utilizes multiple alternative carbon sources simultaneously within a macrophage phagosome, such that these pathways are partially redundant in supporting both energy generation and the substantial biomass addition necessary for hyphal growth and subsequent escape.

We next evaluated whether the alternative carbon mutants would show compromised fitness in the macrophage model, assessing both fungal and macrophage survival after a 16-h coincubation (Fig. 7A and B). In alignment with the hyphal and phagosome acidity phenotypes, the levels of fitness of the alternative carbon mutants were found to be increasingly attenuated as more carbon pathways were eliminated, with the quadruple mutant the most severely impaired. This again suggests that C. albicans utilizes many different nutrient sources upon phagocytosis, a departure from the general paradigm of carbon metabolism in fungi.

Finally, we assessed the virulence of the single and quadruple mutant strains in the mouse intravenous model of disseminated candidiasis. In this model, the virulence of the quadruple mutant was the most severely attenuated compared to the parental strain, and this difference was significantly different from the results seen with the stp2Δ and h-dΔ strains. In this model, the virulence of the jen1Δ jen2Δ strain was slightly attenuated relative to the wild-type strain (the statistical significance was probably the result of the presence of a single outlier that survived much longer than the rest of the members of that experimental group). This contrasts with the macrophage model, in which the jen1Δ jen2Δ strain was the least affected, again pointing to niche-specific differences in carbon availability and importance. Altogether, these data show that C. albicans actively utilizes multiple carbon sources during disseminated infection, which changes fungal morphology, alters stress resistance and immune interactions, and ultimately is essential for pathogenesis.

DISCUSSION

Several lines of evidence demonstrate that the metabolism of nonglucose carbon sources is critical during interactions of C. albicans with phagocytes and in animal models. Many transcriptomic data sets confirmed the occurrence of a wholesale metabolic change upon phagocytosis, with the cell switching from glycolysis to gluconeogenesis (18, 20, 52). This response is not specific to Candida, as many other fungal pathogens, including Cryptococcus neoformans, Aspergillus fumigatus, Talaromyces marneffei, and Paracoccidioides brasiliensis also adapt their metabolism to gain advantages over the host (53–56). Gluconeogenic genes are also induced in vivo (21, 57), and these transcriptional changes are often advantageous for microbial pathogens. Stress resistance induced in the presence of alternative carbon sources has been observed in other Candida species, including C. glabrata and the emerging pathogen C. auris (58, 59), and numerous examples of beneficial effects of utilization of alternative carbon sources have been seen in bacterial pathogens. For example, Staphylococcus aureus, a facultative intracellular pathogen, alters its central carbon metabolism to promote its survival in the phagosome (60, 61). The ability to utilize short-chain fatty acids is required for full virulence of Campylobacter jejuni (62), and Mycobacterium tuberculosis requires the glyoxylate shunt for survival in macrophages and mice (63, 64). Here, we
show that, similarly to many other pathogens, C. albicans appropriately responds to multiple alternative carbon sources to gain advantages over the host.

Mutants impaired in central carbon catabolism via the tricarboxylic acid (TCA)/glyoxylate cycle (icl1Δ strain) or via gluconeogenesis (pck1Δ and fbp1Δ strains) have also been found to be attenuated in macrophage and mouse models (21, 23). Under-
standing which specific nutrients are most relevant in vivo may identify candidate drug targets, as some secondary carbon metabolic pathways, such as the glyoxylate cycle pathway and some amino acid biosynthetic pathways, do not exist or are substantially divergent in mammals. To this end, mutants were previously generated in specific uptake or catabolic pathways, such as those for fatty acids (strain fox2Δ), amino acids (strain stp2Δ), GlcNAC (strain h-dΔ), oligopeptides (strain optΔ ptrΔ), or carboxylic acids (strain jen1Δ jen2Δ), and these have modest or no in vivo phenotypes (19, 23, 32, 33, 65, 66). In contrast, the attenuation of strain ΔΔΔΔ (stp2Δ jen1Δ jen2Δ h-dΔ) in macrophage and mouse models affirms that alternative carbon utilization directly contributes to fungal pathogenesis, and the additive effects observed suggest that C. albicans utilizes multiple carbon sources simultaneously within the macrophage phagosome and during disseminated infection, though we cannot exclude the possibility that the observed differences in virulence resulted in part from other changes, such as changes to the cell wall, induced by variations in nutrient availability.

In addition to the energy generation that is provided by utilizing alternative carbon sources, carboxylic acids, amino acids, and GlcNAC serve as unique signals for C. albicans, modulating stress resistance and macrophage interactions. This was first recognized with lactate, where drug and stress resistance correlated with alterations in the cell wall (25, 27, 28). Amino acids and GlcNAC also induce cell wall alterations and promote stress resistance. Surprisingly, each carbon source confers a unique pattern of resistance; amino acids confer resistance to oxidative and nitrosative stresses, lactate confers resistance to cell wall and osmotic stresses, and GlcNAC confers resistance to oxidative stress. Transcriptomic data provide some basis for these patterns; amino acid-grown cells upregulate several oxidative and nitrosative stress resistance genes, including SOD4 and YHB1 (10), and SOD5 is upregulated by GlcNAC (33). Antifungal resistance to fluconazole is also enhanced by all three classes of compounds and to caspofungin by GlcNAC and amino acids. The abundance of these nutrients in the host differs from site to site, as does the spectrum of immune cells and microbial competition (22). This raises the possibility that C. albicans uses nutrient availability as a signal of its anatomical niche, priming the cell for different arrays of encounters depending on localization.

Indeed, C. albicans is equipped with sensors for all three alternative carbon pathways. Lactate sensing with Gpr1 is required for β-glucan masking and immune evasion, while import and catabolism are dispensable (26). The plasma membrane SPS amino acid sensor (Ssy1, Ptr3, Ssy5) regulates Stp2 and has very similar macrophage phenotypes (67), but it is not clear whether sensing and metabolism can be decoupled. GlcNAC sensing (by Ngs1) and import (through Ngt1) are required for hyphal induction (29), suggesting that catabolism may also be dispensable for GlcNAC-induced stress resistance. Separating nutrient sensing and catabolism for pathogenesis and stress resistance will provide new insights into how pathogenic fungi detect, interact with, and adapt to various host environments.

In addition to the carbon sources studied here, there are likely several other carbon sources that promote fungal survival within the host. β-Oxidation is highly upregulated within macrophages, suggesting that fatty acids such as oleic acid are utilized within the host, although disruption of β-oxidation (fox2Δ) has a minor impact on pathogenesis (23, 24, 68). We mainly focused on the carboxylic acid lactate in this study on the basis of precedents, but preliminary data suggest that C. albicans responds differentially to additional host-relevant carboxylic acids such as α-ketoglutarate and acetate (15, 25, 27). Host-associated sugars besides GlcNAC (sialic acid, galactosamine), in addition to other lipids (e.g., sphingolipids), may be recognized as well.

In conclusion, this report highlights the importance of nutrient flexibility for fungal pathogenesis. In addition to serving as an energy source, alternative carbon sources prime C. albicans to tolerate challenging host environments, ultimately allowing the fungal pathogen to inhabit, survive in, and effectively infect a variety of host niches. Together, these host-relevant carbon sources serve as both an energy source and a
### MATERIALS AND METHODS

**Strains and media.** All strains used in this study are listed in Table 1. For general growth and propagation, *C. albicans* strains were grown in yeast extract-peptone-dextrose (YPD) medium (1% yeast extract, 2% peptone, 2% glucose, 2% agar for solid medium) at 30°C. YPD plus 200 μg/mL nourseothricin (Werner Bioagents, Jena, Germany) was used to select for deletion mutants, and YP maltose (YPM) induced flipping out of the nourseothricin cassette as described previously (69).

Alternative carbon growth and pH experiments required the use of YNB medium with allantoin as the nitrogen source (0.17% yeast nitrogen base without ammonium sulfate and amino acids, 0.5% allantoin) and 1% of the indicated carbon source, adjusted to the indicated pH using HCl or NaOH, similarly to previous publications (10, 13). For the electron microscopy and stress resistance experiments, the medium was buffered to pH 5.5 with 25 mM HEPES.

*J774A.1* murine macrophages were maintained in Dulbecco's modified Eagle's medium (DMEM) plus glutamate and 10% fetal bovine serum (FBS), supplemented with penicillin and streptomycin, at 37°C in a 5% CO₂, humidified environment. DMEM without a pH indicator, penicillin/streptomycin (pen/strep), or FBS was used for the coculture assays.

**Strain construction.** All genetic deletions were generated using the CRISPR/Cas9 SAT-Flipper method (70, 71). Mutants were verified via PCR, and the maltose-inducible SAT cassette was flipped out in order to generate subsequent deletions in the same strain.

Auxotrophic screen mutants from the Noble and Homann libraries were transformed via electroporation with the Cipl30 plasmid digested with Stul (40). All mutants were PCR verified to confirm appropriate gene deletion.

**Growth and pH neutralization assays.** Growth and pH changes were assayed as described previously (10, 14). Briefly, strains were grown overnight in YPD medium at 30°C, centrifuged, washed with YNB media, and diluted into the YNB media for a starting OD₆₀₀ of 0.2. Samples were incubated with aeration for 24 h at 37°C, with growth (OD₆₀₀) and pH assayed every 2 h. Assays were performed in at least triplicate.

**Transmission electron microscopy.** *C. albicans* was grown to mid-log phase in YNB medium plus the indicated carbon source in 50 mL-flasks with shaking at 37°C and then pelleted by low-speed centrifugation, fixed overnight in Karnovsky's fixative, and stored at 4°C until processing for electron microscopy was performed (72). Cells were postfixed in osmium tetroxide, dehydrated in a graded series of ethanol, and embedded in epoxy resin as described previously (73). Sections (100-nm thick) were stained with uranyl acetate and lead citrate. Electron micrographs were collected from randomly selected yeast cells with a JEOL JEM-1230 transmission electron microscope equipped with a digital charge-coupled-device (CCD) camera. All measurements were quantified with ImageJ. The thickness of the cell wall was determined by averaging at least 15 measurements around the wall of each cell, and at least 50 cells were analyzed per condition. Unpaired Student's t tests were performed to determine statistical differences for all measurements.

| Strain | Description | Complete genotype | Reference or source |
|--------|-------------|--------------------|---------------------|
| SC5314 | Wild type   | Prototroph         | 75                  |
| SN250  | Library control + Cipl30 | his1Δ::hisG/his1Δ::hisD leu2Δ::CdHi51/leu2Δ::CMLEU2 arg4Δ::hisG/arg4Δ::hisG | This study |
| MG01   | Constitutive scarlet | al11Δ::LEU2 his1/leu2 arg4/arg4 | This study |
| RW16   | al11Δ + Cipl30 | al11Δ::His1/leu2 arg4/arg4 | This study |
| RW17   | cox4Δ + Cipl30 | cox4Δ::His1/leu2 arg4/arg4 | This study |
| RW18   | oar1Δ + Cipl30 | oar1Δ::His1/leu2 arg4/arg4 | This study |
| RW19   | sin3ΔΔ + Cipl30 | sin3Δ::His1/leu2 arg4/arg4 | This study |
| C_env14| pex13Δ + Cipl30 | pex13Δ::His1/leu2 arg4/arg4 | This study |
| RBV20  | cph1Δ + Cipl30 | cph1Δ::His1/leu2 arg4/arg4 | This study |
| RBV31  | his1/leu2 arg4/arg4 | his1/leu2 arg4/arg4 | This study |
| RW21   | his1/leu2 arg4/arg4 | his1/leu2 arg4/arg4 | This study |
| RW23   | pep8Δ + Cipl30 | pep8Δ::His1/leu2 arg4/arg4 | This study |
| RW24   | swi6Δ + Cipl30 | swi6Δ::His1/leu2 arg4/arg4 | This study |
| RW05   | jen1Δ jen2Δ | jen1Δ::FRT/jen1Δ::FRT jen2Δ::FRT/jen2Δ::FRT | This study |
| RW10   | step2Δ | step2Δ::FRT/step2Δ::FRT | This study |
| RW11   | hxx1Δ nag1 Δdac1Δ | hxx1Δ nag1 Δdac1Δ::FRT/htk1Δ nag1 Δdac1Δ::FRT | This study |
| RW12   | step2Δ jen1Δ jen2Δ | step2Δ::FRT/step2Δ::FRT/step2Δ::FRT | This study |
| RW13   | step2Δ hxx1Δ nag1 Δdac1Δ | step2Δ::FRT/step2Δ::FRT/step2Δ::FRT/htk1Δ nag1 Δdac1Δ::FRT | This study |
| RW14   | jen1Δ jen2Δ hxx1Δ nag1 Δdac1Δ | jen1Δ::FRT/jen1Δ::FRT jen2Δ::FRT/jen2Δ::FRT/htk1Δ nag1 Δdac1Δ::FRT | This study |
| RW15   | jen1Δ jen2Δ step2Δ hxx1Δ nag1 Δ | jen1Δ::FRT/jen1Δ::FRT jen2Δ::FRT/step2Δ::FRT | This study |

*Strains RW16-23 and C_env14 are prototrophic derivatives from the mutant library developed by Noble et al. (40).*
**Stress resistance assays.** *C. albicans* was pregrown for 6 h in the indicated medium with rolling at 30°C and then transferred to a 96-well plate at a starting OD<sub>600</sub> of 0.1. Growth in the presence or absence of the indicated stressor was assayed via OD<sub>600</sub> determination every 15 min using a Cytation 5 plate reader (BioTek) with orbital shaking at 37°C. Growth differences were quantitated as ratios of the changes in OD<sub>600</sub>, after 8 h under the stress conditions to that in the no-stress control (percentage of no stress = ΔOD<sub>600(stress)/ΔOD<sub>600(control) * 100). Assays were performed in at least triplicate, and data were compared using unpaired t tests.

**Macrophage assays.** J774A.1 macrophages were counted using an automated cytometer (Countess II; Thermo Fisher) and seeded at 2.5 × 10<sup>5</sup> cells/ml 24 h prior to initiation of the coculture in DMEM plus glutamate plus FBS plus pen/strep. The medium was replaced with DMEM without FBS, phenol red, or pen/strep immediately prior to the experiment. *C. albicans* strains were subcultured in YPD medium for 4 to 6 h, washed 3 times in phosphate-buffered saline (PBS), counted using the automated cytometer, and added to the macrophages at the indicated multiplicity of infection (MOI). In some experiments, the fungal cells were stained with fluorescein isothiocyanate (FITC) prior to the coculture.

**Germination within macrophages.** FITC-stained *C. albicans* were coincubated with macrophages in phenol red-free DMEM in 8 chamber slides (Ibidi) at an MOI of 1:1 for 2 h. Subsequently, the medium was aspirated and cocultures were fixed with 2.7% paraformaldehyde, counterstained with calceinfluor white (CFW), and maintained in PBS. Bright-field microscopy and fluorescence microscopy were performed with an Olympus IX81 microscope, and germ tube lengths were measured using SlideBook 6 software. Experiments were performed in triplicate, with >30 cells measured per strain. Unpaired Student’s t tests were performed using Prism 8.

**Estimation of phagosome acidity.** Phagosomal pH was estimated using LysoTracker Red, as described previously (11). Briefly, *C. albicans* cells stained with FITC (MOI 1:1) with macrophages preloaded with the acidophilic dye Lysotracker Red for 1 h in phenol red-free DMEM. Samples were fixed with 2.7% paraformaldehyde. Projections of Z-stack fluorescent images were generated to quantify the red fluorescent intensity of 20 total pixels (10 pixels per side of the fungal cell) at points immediately external to the phagocytosed *C. albicans* cells. Values representing the area under the curve (AUC) for 20 pixels per cell are reflective of phagosome acidity. A total of >20 internalized cells were measured per strain. Experiment performed in triplicate. Unpaired Student’s t tests were performed for AUC values in Prism 8.

**Macrophage association assay.** Scarlet fluorescent *C. albicans* cells were pregrown as described above and then coincubated with J774A.1 macrophages stained with NucBlue and ActinGreen under optimal conditions (37°C, 5% CO<sub>2</sub>) in an imaging plate reader (Cytation 5; BioTek). Cocultures were briefly centrifuged to synchronize phagocytosis under each set of conditions. Macrophage associations were automatically calculated over several fields of view by quantifying the number of red *C. albicans* cells associated with the green macrophages (Gen5 software). NucBlue was used for autofocus purposes. Unpaired Student’s t tests were used for statistical analysis.

**Phagocytosis assay.** Scarlet fluorescent *C. albicans* cells were pregrown in alternative carbon sources, coincubated with J774A.1 macrophages stained with NucBlue for 60 min, and then fixed with 2.7% paraformaldehyde and counterstained with (membrane-impermeant) FITC. Several fields of view were imaged using the Cytation 5 plate reader, and phagocytosis rates were calculated automatically by the use of Gen5 software with the following formula: ([total number of red cells] - [number of red fluorescent cells plus number of green fluorescent cells]/[total number of fluorescent red cells]) × 100.

**Fungal survival assay.** Fungal survival during coincubation with macrophages was measured as previously described (74). Macrophages were seeded at 2.5 × 10<sup>5</sup> cells/ml in a 96-well plate 24 h prior to cocubation. *C. albicans* strains were added at 1 × 10<sup>5</sup> cells/ml in wells with or without macrophages, followed by a series of six 1:5 serial dilutions. Microcolonies of *C. albicans* were counted using an inverted microscope in the wells with a discernible number of colonies. Results are represented as percentages of fungal colonies surviving coculture (number of colonies in the presence of macrophages/number of colonies without macrophages × 100) (n = 6). Unpaired Student’s t tests were performed in Prism 8.

**Macrophage survival assay.** Macrophage survival was measured via detection of lactate dehydrogenase (LDH) in culture supernatants using the CytoTox96 nonradioactive cytotoxicity assay (Promega). Log-phase *C. albicans* cells were added at an MOI of 1:1 with macrophages in a 96-well plate and coincubated overnight in phenol red-free DMEM. LDH released by infected macrophages was quantified as described by the manufacturer, corrected for spontaneous LDH release, and compared to the data determined for chemically lysed uninfected macrophages (n = 6). Unpaired Student’s t tests were performed in Prism 8.

**In vivo disseminated candidiasis.** Disseminated *C. albicans* infection was performed as previously described (23). Cells were grown to mid-log phase overnight, centrifuged, washed, counted, and resuspended in PBS. A total of 10 female 7-to-9-week-old ICR mice per strain were infected via tail vein injection with 5 × 10<sup>5</sup> cells/ml in 100 μl PBS for a final concentration of 5 × 10<sup>5</sup> injected cells. Mice were monitored 3 times daily for signs of infection and euthanized when moribund. All animal experiments were conducted according to protocols approved by the Animal Welfare Committee of the University of Texas Health Science Center at Houston. The Mantel-Cox test was performed to statistically compare survival curves.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, PDF file, 0.6 MB.

**FIG S2**, PDF file, 0.05 MB.
REFERENCES

1. Diekema D, Arbeefv依, S, Boyken L, Kroeger J, Pfaller M. 2012. The changing epidemiology of healthcare-associated candidemia over three decades. Diagn Microbiol Infect Dis 73:45–48. https://doi.org/10.1016/j.diagmicrobio.2012.02.001.

2. Pfaller MA, Diekema DJ. 2007. Epidemiology of invasive candidiasis: a persistent public health problem. Clin Microbiol Rev 20:133–163. https://doi.org/10.1128/CMR.00029-06.

3. Concia E, Azizzi AM, Conti M. 2009. Epidemiology, incidence and risk factors for invasive candidiasis in high-risk patients. Drugs 69(Suppl 1):S5–14. https://doi.org/10.2165/11155000-00000000-0000.

4. Cheng SJ, Stoosn LB, Kullberg BJ, Netea MG. 2012. Interplay between Candida albicans and the mammalian innate host defense. Infect Immun 80:1304–1313. https://doi.org/10.1128/IAI.00146-11.

5. Weiss G, Schible UE. 2015. Macrophage defense mechanisms against intracellular bacteria. Immunol Rev 264:182–203. https://doi.org/10.1111/imr.12266.

6. Nathan CF, Hibbs JB. 1991. Role of nitric oxide synthesis in macrophage antimicrobial activity. Curr Opin Immunol 3:65–70. https://doi.org/10.1016/0952-7915(91)90079-g.

7. Fang FC. 2004. Antimicrobial reactive oxygen and nitrogen species: concepts and controversies. Nat Rev Microbiol 2:820–832. https://doi.org/10.1038/nrmicro1004.

8. Flanagan RS, Cosio G, Grinstein S. 2009. Antimicrobial mechanisms of phagocytes and bacterial evasion strategies. Nat Rev Microbiol 7:355–366. https://doi.org/10.1038/nrmicro2128.

9. Kolaiczewska E, Kubes P. 2013. Neutrophil recruitment and function in health and inflammation. Nat Rev Immunol 13:159–175. https://doi.org/10.1038/ni.m3399.

10. Danhof HA, Vykova S, Vesely EM, Ford AE, Gonzalez-Garay M, Lorenz MC. 2016. Robust extracellular pH modulation by Candida albicans during growth in carboxylic acids. mBio 7:e01646-16. https://doi.org/10.1128/mbio.01646-16.

11. Vykova S, Lorenz MC. 26 January 2017, posting date. Phagosomal pH modulation by extracellular acidification is not required for macrophage interleukin 1β production. mBio 8:e00343-12. https://doi.org/10.1128/mbio.00343-12.

12. Lorenz MC, Bender JA, Fink GR. 2004. Transcriptional response of Candida albicans upon internalization by macrophages. Eukaryot Cell 3:1076–1087. https://doi.org/10.1038/35083594.

13. Tucey TM, Verma J, Harrison PF, Ngle ak SL, Lo TL, Scherer AK, Barughare PA, Powell DR, Wheeler RT, Hickey MJ, Beilharz TH, Naderer T, Taven R. 2018. Glucose homeostasis is important for immune cell viability during Candida challenge and host survival of systemic fungal infection. Cell Metab 27:988–1006.e7. https://doi.org/10.1016/j.cmet .2018.03.019.

14. Barelle CJ, Priest CL, MacCallum DM, Gow NA, Odds FC, Brown A. 2006. Niche-specific regulation of central metabolic pathways in a fungal pathogen. Cell Microbiol 8:961–971. https://doi.org/10.1111/j.1462-5822 .2005.00676.x.

15. Miramón P, Lorenz MC. 2017. A feast for Candida: metabolic plasticity confers an edge for virulence. PLoS Pathog 13:e1006144. https://doi.org/10.1371/journal.ppat.1006144.

16. Ramírez MA, Lorenz MC. 2007. Mutations in alternative carbon utilization pathways in Candida albicans attenuate virulence and confer pleiotropic phenotypes. Eukaryot Cell 6:280–290. https://doi.org/10.1111/j.1474-936X.2006.00722.x.

17. Piekaraska K, Mol E, Van Den Berg M, Hardy G, Van Den Burg J, Van Roermund C, MacCallum D, Odds F, Distel B. 2006. Peroxisomal fatty acid β-oxidation is not essential for virulence of Candida albicans. Eukaryot Cell 5:1847–1856. https://doi.org/10.1128/EC.00093-06.

18. Ene IV, Walker LA, Schiavone M, Lee KK, Martin-Yken H, Dague E, Gow NA, Munro CA, Brown A. 2015. Cell wall remodelling enzymes modulate fungal cell wall elasticity and osmotic stress resistance. mBio 6:e00986. https://doi.org/10.1128/mbio.00986-15.

19. Ballou ER, Avelar GM, Childers DS, Mackie J, Bain JM, Wagenier J, Kastora SL, Panea MD, Hardison SE, Walker LA, Erwig LP, Munro CA, Gow NA, Brown GD, MacCallum DM, Brown A. 2016. Lactate signalling regulates fungal β-glucan masking and immune evasion. Nat Microbiol 2:16238. https://doi.org/10.1038/nm microbiol.2016.238.

20. Ene IV, Cheng S, Netea MG, Brown A. 2013. Growth of Candida albicans cells on the physiologically relevant carbon source lactate affects their recognition and phagocytosis by immune cells. Infect Immun 81:238–248. https://doi.org/10.1128/IAI.00109-12.

21. Ene IV, Heilmann CJ, Sorge AG, Walker LA, de Koster CG, Munro CA, Klos FM, Brown A. 2012. Carbon source-induced reprogramming of the cell wall proteome and secretome modulates the adherence and drug resistance of the fungal pathogen Candida albicans. Proteomics 12:3164–3179. https://doi.org/10.1002/pmic.201200228.

22. Naseem S, Min K, Spitzer D, Gardin J, Konopka JB. 2017. Regulation of hyphal growth and N-acetylglucosamine catabolism by two transcription factors in Candida albicans. Genetics 206:299–314. https://doi.org/10.1393/genes/117.101.201491.

23. Sherrington SL, Sorsby E, Mahoney N, Kumwenda P, Lenardon MD, Brown I, Ballou ER, MacCallum DM, Hall RA. 2017. Adaptation of Candida albicans to environmental pH induces cell wall remodelling and enhances innate immune recognition. PLoS Pathog 13:e1006403. https://doi.org/10.1371/journal.ppat.1006403.

24. Alves R, Mota S, Silva S, F Rodrigues C, Alistair AJ, Henriques M, Casal M, Vieira N, Casal M, Johansson B, MacCallum DM, Brown AJP, Paiva S. 2010. The carboxylic acid transporters Jen1 and Jen2 affect the architecture and fluconazole susceptibility of Candida albicans biofilm in the presence of lactate. Biofouling 26:943–954. https://doi.org/10.1080/08927014.2017.1392514.
Functional specialization and differential regulation of short-chain carboxylic acid transporters in the pathogen Candida albicans. Mol Microbiol 75:1337–1354. https://doi.org/10.1111/j.1365-2958.2009.07003.x.

33. Naseem S, Araya E, Konopka JB. 2015. N-Acetylglucosamine regulates virulence properties in Candida albicans. PLoS Pathog 11:e1004947. https://doi.org/10.1371/journal.ppat.1004947.

34. Westman J, Moran G, Mogavero S, Hube B, Grinstein S. 2018. Candida albicans hyphal expansion causes phagosomal membrane damage and hypoosmotic alkalization. mBio 9:00122e-18. https://doi.org/10.1128/mBio.00122-18.

35. Fradin C, De Groot P, MacCallum D, Schaller M, Kils F, Odds FC, Hube B. 2005. Granulocytes govern the transcriptional response, morphology and proliferation of Candida albicans in human blood. Mol Microbiol 56:397–415. https://doi.org/10.1111/j.1365-2958.2005.04575.x.

36. Morschhäuser J. 2013. Roles of different peptide transporters in nutrient acquisition and immune evasion in Candida albicans. Microbiology 159:963–973. https://doi.org/10.1099/micro.0.06178-0.

37. Lopes-Oliveira F, Soares-da-Silva P, Almeida-Sousa CA, da Cunha Pinheiro LT, Sá-Lopes I. 2010. Transcriptional induction of a galactose responsive ABC transporter in the pathogen Candida albicans. Mol Microbiol 75:105–118. https://doi.org/10.1111/j.1365-2958.2009.05869.x.

38. Kji H, Kim M, Cho H, Lee S, Kim Y, Park Y, Shi J, Kim J, Kim H, Kim J. 2015. A role of Pex5p in respiration metabolism of T. maritima. J Microbiol Biotechnol 25:188–194. https://doi.org/10.4014/jmb.1411.11001.

39. Reuvers PA, Groen JF, Staal TP, de Boer AJ, de Vos WM, de Groot J, Reemtsma K. 2015. Role of snRNP Pathway during malate synthase inactivation in the pathogen Candida albicans. Microbiology 161:1731–1742. https://doi.org/10.1099/micro.0.08614-0.

40. Da Fonseca G, Silva L, Pinto L, Santos M, Cunha A, Leite M, Costa A, Cunha I. 2003. Expression of iron-regulated ABC transporter genes and transcriptional response during different iron-limited conditions in the pathogen Candida albicans. Gene 312:109–119. https://doi.org/10.1016/S0378-1119(02)01664-8.

41. Mardor G, Bruschi S, da Silva W, Zhou K, Li C, Liu Y. 2015. The yeast orthologue of OPII. Yeast 32:793–907. https://doi.org/10.1002/yea.3104.

42. Fan W, Kraus PR, Boily MJ, Heitman J. 2015. Cryptococcus neoformans genic expression during murine macrophage infection. Eukaryot Cell 14:2140–1333. https://doi.org/10.1128/EC.01430-1335.

43. Turcotte B, Liang XB, Robert F, Soontorngun N. 2010. Transcriptional coregulation by the cell integrity mitogen-activated protein kinase Slt2 in C. albicans. Mol Microbiol 77:495–507. https://doi.org/10.1111/j.1365-2958.2010.06590.x.

44. Ramírez-Zavala B, Mottola A, Haubenreißer J, Schneider S, Allert S, Tempst P, Sabatini DM. 2002. mTOR interacts with raptor to form a heterodimeric protein kinase complex. EMBO J 21:4577–4587. https://doi.org/10.1093/emboj/cdh475.

45. Fan W, Kraus PR, Boily MJ, Heitman J. 2015. Cryptococcus neoformans genic expression during murine macrophage infection. Eukaryot Cell 14:2140–1333. https://doi.org/10.1128/EC.01430-1335.

46. Ramírez MA, Lorenz MC. 2009. The transcription factor homolog CTF1 boxylic acid transporters in the pathogen Candida albicans. Mol Microbiol 75:1337–1354. https://doi.org/10.1111/j.1365-2958.2009.07003.x.

47. Heyken WT, Wagner C, Wittmann J, Albrecht A, Schüller HJ. 2003. Analysis of the regulation, expression, and localisation of the isocitrate lyase from C. albicans. Eukaryot Cell 2:118–129. https://doi.org/10.1128/EC.00206-09.

48. Chan WT, Swenson D, Sacchettini JC, Jacobs WR, Russell DG. 2000. Intracellular pathogen coregulation by the cell integrity mitogen-activated protein kinase Slt2 in C. albicans. Mol Microbiol 36:393–405. https://doi.org/10.1046/j.1365-2958.2000.01847.x.

49. Fantes P, Nurse P. 1977. Control of cell size at division in fission yeast by a growth-modulated size control over nuclear division. Exp Cell Res 107:377–386. https://doi.org/10.1016/0014-4827(79)90359-7.

50. Johnston GC, Pringle JR, Hartwell LH. 1977. Coordination of growth with cell division in the yeast Saccharomyces cerevisiae. Exp Cell Res 105: 79–98. https://doi.org/10.1016/0014-4827(79)90359-7.

51. Kim DH, Sarbasov DA, Ali SM, King JE, Latek RR, Erdjument-Bromage H, Tempst P, Saldanha JP. 2002. mTOR cooperates with raptor to form a tumor suppressor complex that signals to the cell division machinery. Cell 110:163–175. https://doi.org/10.1016/S0092-8674(02)00808-5.

52. Briscoe S, Cooper CR, Vanittanakom N. 2008. Molecular analysis of the Penicillium marneffei glyceraldehyde-3-phosphate dehydrogenase orthologue of OPII. Yeast 20:1177–1188. https://doi.org/10.1002/yea.1031.

53. Turcotte B, Liang XB, Robert F, Soontorngun N. 2010. Transcriptional coregulation by the cell integrity mitogen-activated protein kinase Slt2 in C. albicans. Mol Microbiol 77:495–507. https://doi.org/10.1111/j.1365-2958.2010.06590.x.

54. Elgersma Y, Kwast L, Klein A, Voorn-Brouwer T, van den Berg M, Metzig C, America T, Tabak HF, Distel B. 1996. The SH3 domain of the Saccharomyces cerevisiae Pex5p receptor mediates interaction with a docking site for Pex5p, a mobile receptor for the import PTS1-containing proteins. J Cell Biol 135:97–109. https://doi.org/10.1083/jcb.135.1.97.

55. Sánchez-Aguilera LF, Cárdenas-Sánchez A, de la Fuente C, Torres-Paredes M, Fernández-Escobar I, Arroyo A, Vargas A, Alcaraz S, Cervera M, Pascual J, Pérez-Salmerón A. 2016. The yeast ortholog of OPII mediates protein import into the yeast mitochondria. Biol Chem 397:367–377. https://doi.org/10.1515/bsl-2015-0108.

56. Thirach S, Cooper CR, Vanittanakom N. 2008. Molecular analysis of the Penicillium marneffei glyceraldehyde-3-phosphate dehydrogenase orthologue of OPII. Yeast 20:1177–1188. https://doi.org/10.1002/yea.1031.
69. Huang MY, Mitchell AP. 2017. Marker recycling in Candida albicans through CRISPR-Cas9-induced marker excision. mSphere 2:e00050-17. [https://doi.org/10.1128/mSphere.00050-17].

70. Vyas VK, Barrasa MI, Fink GR. 3 April 2015, posting date. A Candida albicans CRISPR system permits genetic engineering of essential genes and gene families. Sci Adv [https://doi.org/10.1126/sciadv.1500248].

71. Min K, Ichikawa Y, Woolford CA, Mitchell AP. 2015. Candida albicans gene deletion with a transient CRISPR-Cas9 system. mSphere 1:e00130-16. [https://doi.org/10.1128/mSphere.00130-16].

72. Karnovsky MJ. 1965. A formaldehyde-glutaraldehyde fixative of high osmolarity for use in electron microscopy. J Cell Biol 27:1A–149A.

73. Baker PA, Meyer MD, Tsang A, Uribe RA. 6 May 2019, posting date. Immunohistochemical and ultrastructural analysis of the maturing larval zebrafish enteric nervous system reveals the formation of a neuropil pattern. Sci Rep [https://doi.org/10.1038/s41598-019-43497-9].

74. Rocha CRC, Schröppel K, Harcus A, Dignard D, Taylor BN, Thomas DY, Whiteway M, Leberer E. 2001. Signaling through adenylyl cyclase is essential for hyphal growth and virulence in the pathogenic fungus Candida albicans. Mol Biol Cell 12:3631–3643. [https://doi.org/10.1091/mbc.12.11.3631].

75. Gillum AM, Tsay EY, Kirsch DR. 1984. Isolation of the Candida albicans gene for orotidine-5'-phosphate decarboxylase by complementation of S. cerevisiae ura3 and E. coli pyrF mutations. Mol Gen Genet 198:179–182. [https://doi.org/10.1007/bf00328721].