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Carnitine-Dependent Transport of Acetyl Coenzyme A in *Candida albicans* Is Essential for Growth on Nonfermentable Carbon Sources and Contributes to Biofilm Formation

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In eukaryotes, acetyl coenzyme A (acetyl-CoA) produced during peroxisomal fatty acid β-oxidation needs to be transported to mitochondria for further metabolism. Two parallel pathways for acetyl-CoA transport have been identified in *Saccharomyces cerevisiae*: one is dependent on peroxisomal citrate synthase (Cit), while the other requires peroxisomal and mitochondrial carnitine acetyltransferase (Cat) activities. Here we show that the human fungal pathogen *Candida albicans* lacks peroxisomal Cit, relying exclusively on Cat activity for transport of acetyl units. Deletion of the *CAT2* gene encoding the major Cat enzyme in *C. albicans* resulted in a strain that had lost both peroxisomal and mitochondrion-associated Cat activities, could not grow on fatty acids or C2 carbon sources (acetate or ethanol), accumulated intracellular acetyl-CoA, and showed greatly reduced fatty acid β-oxidation activity. The cat2 null mutant was, however, not attenuated in virulence in a mouse model of systemic candidiasis. These observations support our previous results showing that peroxisomal fatty acid β-oxidation activity is not essential for *C. albicans* virulence. Biofilm formation by the cat2 mutant on glucose was slightly reduced compared to that by the wild type, although both strains grew at the same rate on this carbon source. Our data show that *C. albicans* has diverged considerably from *S. cerevisiae* with respect to the mechanism of intracellular acetyl-CoA transport and imply that carnitine dependence may be an important trait of this human fungal pathogen.

β-Oxidation of fatty acids is a well-conserved metabolic process that results in the stepwise degradation of fatty acids to acetyl coenzyme A (acetyl-CoA). While this process takes place both in mitochondria and peroxisomes in mammalian cells, oxidation of fatty acids is exclusively peroxisomal in plant and fungal cells (16). To allow further metabolism, the acetyl-CoA produced must be transported from peroxisomes to the mitochondria and fed into the mitochondrial TCA cycle and is oxidized completely to CO2 and H2O. In the yeast *Saccharomyces cerevisiae*, two pathways for the transport of acetyl-CoA from peroxisomes to mitochondria have been identified; the first pathway depends on peroxisomal citrate synthase (Cit2p), while the second pathway requires carnitine acetyltransferase (Cat2p) (40). In the Cat2p-dependent pathway, the acetyl moiety of peroxisomal acetyl-CoA is linked to oxaloacetate by Cit2p to form citrate and free CoA. Citrate is subsequently transported to the mitochondria, where it enters the TCA cycle. In the Cat2p-dependent pathway, Cat2p catalyzes the formation of acetyl-carnitine from acetyl-CoA and the carrier molecule carnitine. Formation of acetyl-carnitine is necessary to allow the transport of acetyl units over the peroxisomal and mitochondrial membranes. Mitochondrial Cat2p catalyzes the reverse reaction, liberating the acetyl unit from carnitine and coupling it to a molecule of free CoA for further metabolism. Interestingly, in *S. cerevisiae*, the peroxisomal and mitochondrial forms of Cat2p are encoded by a single gene and the mechanism of differential targeting of the protein has been well established (6). The work of van Roermund et al. (39, 40) has shown that the Cit2p- and Cat2p-dependent pathways of acetyl-CoA export can work in parallel; disruption of either the *CIT2* or the *CAT2* gene in *S. cerevisiae* does not lead to a growth defect on fatty acids, while the simultaneous disruption of both genes does. While both pathways function in *S. cerevisiae*, plants seem to lack the Cat pathway, as disruption of both peroxisomal citrate synthases (CSY2 and CSY3) in *Arabidopsis thaliana* resulted in a mutant that failed to metabolize triacylglycerol (27).

In *S. cerevisiae*, acetyl-CoA transport can be studied independently of fatty acid β-oxidation. When yeast cells are growing on acetate or ethanol, acetyl-CoA is formed in the cytosol, which needs to be transported to the mitochondria and fed into the glyoxylate cycle. Transport of acetyl-CoA from the cytosol to mitochondria is probably mediated by one or both of the other carnitine acetyltransferases in *S. cerevisiae*, Yat1p and Yat2p. Yat1p is localized to the outer mitochondrial membrane (33), while the localization of Yat2p is unknown. The
specific functions of either of the Yat proteins are not known; however, a random-mutagenesis screen of an *S. cerevisiae* *cit2Δ* strain (which is dependent on the Cat pathway for growth on oleate and ethanol) revealed that all three carnitine acyltransferases (Cat2p, Yat1p, and Yat2p) are essential for growth on nonfermentable carbon sources such as acetate, ethanol, oleate, and glycerol (35).

The *C. albicans* genome encodes three putative carnitine acyltransferases, homologs of *S. cerevisiae* Cat2p, Yat1p, and Yat2p, which have been named Ctn2p, Ctn1p, and Ctn3p, respectively (28). Mitochondrial and peroxisomal targeting signals are conserved in the *C. albicans* Cat2p homolog (Ctn2p), and two in-frame start codons are present, as is also the case in *S. cerevisiae* (6), suggesting that the mechanism of dual localization of Cat2p is conserved between the two yeasts. All three putative carnitine acyltransferases are up-regulated during phagocytosis by macrophages (28).

To allow growth on fatty acids, ethanol, or acetate, not only must the acetyl-CoA produced be transported to the mitochondria for oxidation in the TCA cycle, but this C2 compound also needs to be converted to C4 units (succinate) that can be used for biosynthetic purposes. The metabolic pathway that allows the net synthesis of C4 units from acetyl-CoA is the glyoxylate cycle (14). The essential role of this metabolic pathway for the utilization of nonfermentable carbon sources has been substantiated through the analysis of fungal mutants lacking one of the key enzymes of the glyoxylate cycle, i.e., isocitrate lyase (Icl1p) or malate synthase (Mls1p). *S. cerevisiae* icel1 or mls1 mutants are unable to grow on fatty acids, ethanol, or acetate as the sole carbon source (7, 11, 21), and similar phenotypes have been reported for *C. albicans* mutants lacking Icl1p (20, 26). While the function of the glyoxylate pathway in eukaryotes is well conserved, the subcellular localization of the enzymes may vary from organism to organism. For example, in plants and certain fungi such as *C. albicans*, Icl1p and Mls1p are exclusively peroxisomal (24; our unpublished data), whereas in *S. cerevisiae*, Icl1p is cytosolic and Mls1p is only peroxisomal when cells are grown on fatty acids (17, 36).

Interestingly, acetyl-CoA transport and metabolism seem to play an essential role in the virulence of pathogenic fungi. The glyoxylate pathway enzymes Icl1p and Mls1p were shown to be crucial for the virulence of the plant pathogens *Magnaporthe grisea* (44) and *Stagonospora nodorum* (34) and the human pathogen *C. albicans* (20, 26), while carnitine-dependent acetyl unit transport in *M. grisea* is required for the elaboration of penetration hyphae during plant infection (2, 30).

In this study, we investigated the roles of acetyl-CoA transport in fatty acid metabolism and in the virulence of the human fungal pathogen *C. albicans*. We demonstrate here that the *C. albicans* genome contains only one citrate synthase (*CIT*) gene (alleles orf19.4393 and orf19.11871) and show that citrate synthase activity is present in mitochondria but not in peroxisomes, suggesting that for growth on fatty acids and transport of acetyl units from peroxisomes, the fungus is dependent on the Cat pathway. A *C. albicans* mutant lacking the major Cat protein (Cat2p) is not able to grow on fatty acids or on acetate or ethanol, but virulence of the mutant is not attenuated. While the cat2 deletion strain exhibits no growth deficiency on glucose, the mutant shows a small but significant reduction in its ability to form biofilms in vitro on this carbon source. Our results provide insight into the mechanism of intracellular acetyl-CoA transport in *C. albicans* and suggest that carnitine dependency may be an important trait of this human fungal pathogen.

**Materials and Methods**

**Media and culture conditions.** *C. albicans* strains were grown at 28°C unless otherwise stated. For routine nonselective culturing of *C. albicans* strains, YPD (2% Bacto peptone, 1% yeast extract, 2% glucose, 20 μg/ml uridine) was used. *C. albicans* transformants were selected and grown on minimal solid medium containing 0.67% yeast nitrogen base (YNB) without amino acids (Difco), 2% glucose, 2% agar, 80 μg/ml uridine, and amino acids as needed (20 μg/ml arginine, 20 μg/ml histidine). Plates used for spot assays had the same composition and contained glucose (2%), oleic acid-Tween 80 (0.12%–0.2%), ethanol (2%), or sodium acetate (2% with 0.5% potassium phosphate buffer [pH 6.0]) as a carbon source. For Cat enzyme assays, strains were grown overnight on YPD, YPO (2% Bacto peptone, 1% yeast extract, 0.12%–0.2% oleic acid-Tween 80), YPA (0.5% Bacto peptone, 0.3% yeast extract, 0.5% potassium phosphate buffer [pH 6], 2% sodium acetate), or YPE (0.5% Bacto peptone, 0.3% yeast extract, 1.5% potassium phosphate buffer [pH 6], 2% ethanol). For subcellular fractionation, β-oxidation assay, and immunoelectron microscopy, strains were grown overnight on YPO. In all experiments, strains were pregrown on minimal glucose medium (0.3% glucose, 0.67% YNB) for at least 8 h before being shifted to the medium of choice.

**Spot tests.** Cells were pregrown on medium containing 0.3% glucose, washed, resuspended to a concentration of about 2.7 × 10^7 cells/ml, and serially diluted (1:10 dilutions). Four microliters of each dilution was spotted onto agar plates. The pictures were taken after 3 days (glucose) or 5 days (oleate, acetate, and ethanol) of incubation at 28°C. Sensitivity to various stress agents was tested by spotting serial dilutions of cells onto YPD plates containing 25 μg/ml Calcofluor white, 200 μg/ml Congo red, 1.5 M sorbitol, or 0.05% sodium dodecyl sulfate. Plates were incubated at 28°C for 3 days. Hypha formation was tested by spotting dilutions onto YPD plates containing 10% fetal calf serum or 3% glycerol, followed by incubation at 37°C for 7 days.

**Strains and plasmids.** The *C. albicans* strains used in this study (listed in Table 1) are derivatives of SN76 (23). The plasmids and primers used in this study are listed in Tables 2 and 3, respectively. The CAT2 gene (alleles orf19.4591 and orf19.12806) was deleted by using a PCR-based procedure with primers containing 100-bp regions identical to the 5′ and 3′ flanking sequences of the open reading frames (45). A cat2Δcat2 Δ strain was created by successive transformation with two disruption cassettes. The first cassette, containing the ARG4 auxotrophic marker, was amplified from plasmid pFA-CaARG4 (8) by using primers CAT2-D-F-FA and CAT2-D-R-FA in combination with extension primers CAT2-F-Ext and CAT2-R-Ext. The same primers were used for construction of the second cassette by amplification on plasmid pFA-ChHS1 (32), which contains the HIS1 auxotrophic marker from *Candida dublinensis*. The cat2Δcat2 Δ strain was transformed with Xhol-digested pLUBP (29) to create a prototrophic cat2 null mutant. For complementation of the cat2Δcat2 Δ strain with the CAT2 gene, a plasmid was constructed by PCR with primers KS128 and JB10 on genomic DNA isolated from SN76. The PCR product, containing the 800-bp promoter region, the full-length CAT2 open reading frame, and the 800-bp 3′ untranslated region, was cloned into pLUBP, resulting in pLUBP pKa30. Transformation of the cat2Δcat2 Δ strain with Xhol-digested pKa30 resulted in the cat2ΔΔ + CAT2 strain (the complemented deletion strain). A prototrophic SN76 strain (SN76-P) was constructed by sequential transformation of strain SN76 with Nhel-linearized pLUBP (for complementation of the *URA3-IR01* region), a 2.0-kb Kpn1/SacI fragment of pRS-ARG4*SpeI* (to complement the ARG4 deletion) and a 1.8-kb BamHI/XbaI fragment from pGEM-HIS1 (to complement the HIS1 deletion). Strain SN76-P will be referred to as the wild type. All strains were verified by PCR.

**Transformation.** *C. albicans* was transformed by using a modified lithium acetate protocol (42). Heat shock was carried out at 44°C for 15 min.

Subcellular fractionation and density gradient analysis of *C. albicans*. Cells were inoculated in YNB–2% glucose and grown overnight. On the next morning, the culture was diluted in YNB–0.3% glucose to an optical density at 600 nm (OD600) of 0.15, grown to an OD600 of 1.0 to 1.5, and then inoculated into YPO at an OD600 of 0.01 and grown overnight. On the next morning, cells were collected by centrifugation (10 min at 4,000 × g), washed three times with 25 ml distilled water, and collected again by centrifugation (10 min at 4,000 × g). Cells were resuspended in 5 ml of buffer Z (0.5 M KCl, 5 mM morpholinepropane-
sulfonic acid [MOPS; pH 7.2], 10 mM Na2SO4) containing 0.25 mg Zymolyase 100T (ICN Biomedicals) per g (wet weight) and incubated for 10 to 30 min at 28°C while shaking at 120 rpm to convert the cells to spheroplasts. Spheroplast formation was monitored microscopically. Spheroplasts were harvested by centrifugation at 2,300 g at 4°C to remove cell debris and nuclei, and the low-speed supernatant (H) was separated into an organellar pellet (P) and a cytosolic (S) fraction by centrifugation for 20 min at 20,000 x g at 4°C. The organellar pellet was taken up in 1 ml buffer G (5 mM Tris-HCl [pH 7.5], 3 mM KCl, 0.3 mM EDTA [pH 8], 0.1% ethanol, 0.6 M sorbitol) and loaded onto a 15 to 50% Nycodenz gradient. Gradients were spun for 5 min at 4°C and the supernatant was transferred to a glass tube and evaporated under a stream of nitrogen at 40°C. The final residue (a mitochondrial marker), and carnitine acetyltransferase (Cat) activity (6). Subcellular fractionation and density gradient analysis of S. cerevisiae were performed as described previously (6).

**Fatty acid β-oxidation measurements.** The β-oxidation activity in intact cells was measured essentially as described before (39), except that the cells were resuspended at an OD600 of 1. Production of CO2 and incorporation of radioactive label into acid-soluble matter were followed over time.

**Acyl-CoA measurements.** Acyl-CoA measurements were performed as described by Hammond et al. (10), with some modifications. Approximately 20 mg of freeze-dried material of oleate-grown yeast cells was added to 1.5 ml Eppendorf vials, and the exact weight of the sample was determined with a microbalance. To the sample, 20 μl of internal standard (2H3-acetyl-CoA [100 μM], 2H3-octanoyl-CoA [100 μM], and 2H3-palmitoyl-CoA [100 μM]) in 70% acetone (nitrile) was subsequently added on ice in 50 mM KH2PO4–50% 2-propanol. An equal volume of acetonitrile was added. After 3 min of mixing, the samples were centrifuged at 1,600 x g for 5 min at 4°C and the supernatant was transferred to a glass tube and evaporated under a stream of nitrogen at 40°C. The final residue was taken up in 100 μl methanol-H2O (1:1) and subjected to liquid chromatography-tandem mass spectrometry analysis as described before (10).

**Virulence studies.** Virulence assays were performed at the University of Aberdeen by using a murine tail vein injection model. Groups of six BALB/c female mice were challenged intravenously with the C. albicans strains at a dose of 1.5 × 104 CFU/g body weight as previously described (26). Mice were weighed and monitored daily for 14 days. Mice that died or were humanely killed were recorded as dead. Surviving mice were considered to be cured at this end point. As a control, untreated inocula of strain C. albicans were conducted in parallel for each experiment.

**Acyl-CoA dehydrogenase activity (43) (a peroxisomal marker), fumarase activity (38) (a mitochondrial marker), and carnitine acetyltransferase (Cat) activity (6).** Subcellular fractionation and density gradient analysis of S. cerevisiae were performed as described previously (6).

### TABLE 2. Plasmids used in this study

| Plasmid | Purpose | Reference |
|---------|---------|-----------|
| pLUBP   | URA3 complementation | 29 |
| pFa-CaARG4 #201 | ARG4 disruption | 8 |
| pFA-CdHIS1 #627 | CdHIS1 disruption | 32 |
| pGEM-HIS1 | HIS1 complementation | 45 |
| pRS-ARG4 | ARG4 complementation | 45 |
| pKa30 | CAT2 complementation | This study |

### TABLE 3. Primers used in this study

| Primer | 5’–3’ sequence |
|--------|----------------|
| CaCAT2-D-F-FA | AACTAATCAGAAGAGATAGGTCGAA |
| CaCAT2-D-R-FA | AAAATTTAAGACCTTTATATGTCAT |
| CaCAT2-Ext-F | GTGCTAATAATAACTAAATTTAAGAA |
| CaCAT2-Ext-R | ATTCGAG |
| CAT2-C-F | ATCAAGTATCATGACCCCCAC |
| CAT2-C-R | GATTGAGATGTGTTGAGG |
| CaA2 | GAGCATCAGTCGACCCCGG |
| X2-CdHIS1 | TCTAAATCAGAAGAGATAGGTCGAA |
| X3-CdHIS1 | GATCAGTCGACCCCGG |
| JB10 | ATGTTGTTACCAAGAGTGTGTTAGG |
| KS128 | TGGAGCCCTCGTTGATTTAAATCCTCGG |

### TABLE 1. Yeast strains used in this study

| Strain | Species | Genotype | Reference |
|--------|---------|----------|-----------|
| BJ1991 | S. cerevisiae | MATa leu2 ura3::251 trp1::pb1-1122 pep4-3 gal2 | 13 |
| SN76 | C. albicans wild-type auxotroph | arg4/arg4 his1::HIS1 ura3::HIS4 trp1::HIS4 | 23 |
| SN76-P | C. albicans wild-type prototroph | ARG4 HIS4 complementation 29 | This study |
| CEM28 | C. albicans | cat2Δ/cat2Δ | This study |
| CEM38 | C. albicans | cat2Δ/cat2Δ + URA3 | This study |
| CKS58 | C. albicans | cat2Δ/cat2Δ + CAT2 | This study |
| CEM16 | C. albicans | fox2Δ/fox2Δ + URA3 | 26 |

| Strain | Species | Name | Genotype | Reference |
|--------|---------|------|----------|-----------|
| SN76   | C. albicans | arg4Δ/arg4Δ his1::HIS1 ura3::HIS4 trp1::HIS4 | This study |
| SN76-P | C. albicans | arg4Δ/ARG4 HIS4 his1::HIS1 ura3::HIS4 trp1::HIS4 | This study |
| CEM28  | C. albicans | cat2Δ/cat2Δ | This study |
| CEM38  | C. albicans | cat2Δ/cat2Δ + URA3 | This study |
| CKS58  | C. albicans | cat2Δ/cat2Δ + CAT2 | This study |
| CEM16  | C. albicans | fox2Δ/fox2Δ + URA3 | This study |
observed daily. Animals that developed signs of serious illness or lost more than 20% of their initial body weight were humanely terminated and recorded as dying the following day. Survival data were analyzed by log rank statistics, and tissue burden data were analyzed by \( t \) test.

Growth and analyses of biofilms. Biofilms were grown in 96-well plates (Costar; Corning Incorporated, Corning, NY) coated with fetal bovine serum (FBS) as described previously (15), and growth was quantified by crystal violet staining (37). Biofilm thicknesses were measured with a low-load compression tester (LLCT) (25) on biofilms grown on FBS-coated (1.5 by 1.5 cm) polymethylmethacrylate slides in 12-well tissue culture plates. For confocal laser scanning microscopy (CLSM), biofilms were grown in FBS-coated 12-well plates, washed once with 2 ml phosphate-buffered saline (10 mM potassium phosphate, 0.15 M NaCl [pH 7.0]), and incubated with 1 ml phosphate-buffered saline containing BacLight (Molecular Probes, Leiden, The Netherlands), prepared according to the manufacturer’s instructions, for 30 min at room temperature in the dark. Confocal laser scanning microscope model LEICA TCS SP2 (Leica Microsystems Heidelberg GmbH, Heidelberg, Germany) equipped with a He-Ne laser and an Ar laser and supplied with the latest version of the Leica Confocal software was used to visualize the submerged biofilms with a 40× water objective. The two components of BacLight were excited at 488 nm, the green signal was recorded with a 510- to 540-nm emission filter setting, and the red signal was recorded with a 600- to 680-nm emission filter setting.

RESULTS

C. albicans contains one citrate synthase that is localized to mitochondria. The S. cerevisiae genome encodes three citrate synthases, of which two are mitochondrial (Cit1p and Cit3p) and one is peroxisomal (Cit2p) (9, 12, 18, 31). We performed BLASTp searches with the S. cerevisiae Cit proteins and found that the C. albicans genome (http://www.candidagenome.org) contains only one gene predicted to encode citrate synthase. The putative C. albicans citrate synthase (orf19.4393) shows 77% sequence identity with ScCit1p and 72% and 45% identity with ScCit3p and ScCit2p, respectively, and has a predicted mitochondrial targeting signal in its N-terminal region (4). To determine the subcellular localization of the C. albicans citrate synthase in the gradients was determined by immunoblotting with an antibody directed against S. cerevisiae citrate synthase 1 (Cit1p).
synthase, we performed subcellular fractionation, followed by density gradient analysis, on cells grown in rich medium containing oleic acid (Fig. 1A). Analysis of the *C. albicans* gradient fractions by immunoblotting with an antibody directed against *S. cerevisiae* Cit1p revealed a single band with an apparent molecular mass of about 45 kDa, corresponding to the predicted molecular mass of the *C. albicans* Cit1p protein that cofractionated with the mitochondrial marker enzyme fumarase. Very little signal was detected in the peroxisomal peak fractions, suggesting that the single CIT gene in *C. albicans* encodes a mitochondrion-targeted protein.

As a control, oleate-grown *S. cerevisiae* cells were analyzed in a similar way. Figure 1B shows that the anti-Cit1 antibody recognizes at least two citrate synthase proteins in *S. cerevisiae*, a band running at about 45 kDa that cofractionates with the mitochondrial marker and most likely represents Cit1p and/or Cit3p and a band enriched in the peroxisomal fractions that runs slightly faster, presumably corresponding to peroxisomal Cit2p. Because of the almost identical predicted molecular masses of Cit1p and Cit3p, it is very possible that both proteins run at the same position in the gel and are observed as a single band or, alternatively, that either of the two proteins is not expressed under our experimental conditions. Notwithstanding the above, these data confirm previous results of Lewin et al. (18), who showed the presence of a peroxisomal citrate synthase (Cit2p) in *S. cerevisiae* and imply that peroxisomes in *C. albicans* lack this protein.

**Carnitine acetyltransferase is abundantly present in peroxisomes and mitochondria of *C. albicans*.** The absence of peroxisomal citrate synthase in *C. albicans* predicts that for export of acetyl units, the organism depends on the carnitine acetyltransferase pathway. We measured Cat activity in the gradient fractions of both *C. albicans* and *S. cerevisiae* (Fig. 1A and B). Cat activity in the *C. albicans* gradient colocализed with the peroxisomal and mitochondrial peaks, and the activities in the two peaks are comparable. In concordance with Elgersma et al. (6), we also found a bimodal distribution of the Cat activity in *S. cerevisiae*, with about five times more activity in the mitochondrial peak fraction than in the peroxisomal peak fraction.

**CAT2 encodes the major carnitine acetyltransferase in *C. albicans*.** In *S. cerevisiae*, the CAT2 gene encodes both the peroxisomal and mitochondrial forms of Cat2p and contributes about 95% of the total Cat activity in oleate-grown cells, while the remaining activity is ascribed to Yat1p and Yat2p (6, 35). To determine the role of Cat2p in *C. albicans*, we constructed a CAT2 (orf19.4591 and orf19.12060) deletion strain by a PCR-based gene disruption procedure (45) (see Materials and Methods). To generate the complemented strain (cat2Δ/cat2Δ + CAT2), the CAT2 gene, together with the 800-bp upstream promoter region, was cloned into the pLUBP vector (29) and transformed into the cat2Δ/cat2Δ strain. URA3 prototrophy was restored to the cat2Δ/cat2Δ strain by introducing the empty pLUBP vector.

Total Cat activity was determined in lysates of the wild-type, cat2Δ/cat2Δ, and complemented strains grown on rich medium containing glucose, oleate-Tween 80, ethanol, or acetate (Table 4). The Cat activity of the wild-type and complemented strains was about twofold higher on oleate, acetate, and ethanol compared to that on glucose. The Cat activity of the cat2Δ/cat2Δ strain was just above the background levels under the growth conditions tested, in spite of the fact that the *C. albicans* genome encodes two other putative carnitine acetyltransferases, potential homologs of *S. cerevisiae* Yat1p and Yat2p (28, 33, 35). We tried to detect residual Cat activity in the cat2Δ/cat2Δ strain by varying the conditions of the assay (dif-

### TABLE 4. Cat activities in lysates of *C. albicans* cells grown on various carbon sources

| Strain                  | Glucose       | Oleate       | Acetate      | Ethanol      |
|-------------------------|---------------|--------------|--------------|--------------|
| Wild type               | 653.4 ± 18.2  | 1,189.5 ± 403.3 | 1,273.8 ± 3.0 | 1,486.2 ± 39.9 |
| cat2Δ/cat2Δ             | 1.1 ± 0.3     | 0.1 ± 0.1    | 3.4 ± 5.5    | 0.9 ± 1.8    |
| cat2Δ/cat2Δ + CAT2      | 517.3 ± 59.3  | 1,403.9 ± 115.4 | 939.5 ± 11.9 | 679.9 ± 21.4 |

* Nanomoles of [14C]-acetylcarnitine formed per minute per milligram of protein.
different pH, addition of a nonionic detergent) by varying the method of protein lysate preparation and by using purified peroxisomal and mitochondrial fractions instead of total protein lysates (data not shown). We were, however, not able to detect significant amounts of Cat activity in the cat2 null mutant by any of these methods, suggesting that the expression of the YAT genes is very low under the conditions tested. Taken together, these results show that the C. albicans CAT2 gene, like the Saccharomyces C. C. gene (6), encodes both the major peroxisomal and mitochondrial forms of Cat.

The C. albicans cat2 null mutant is unable to utilize fatty acids, ethanol, or acetate. To determine whether, in the absence of a peroxisomal citrate synthase, peroxisomal Cat2p plays an essential role in the export of the acetyl-CoA produced during fatty acid β-oxidation, we tested the cat2Δ/cat2Δ strain for the ability to utilize oleate as the sole carbon source. Serial dilutions of the wild-type, cat2Δ/cat2Δ, and complemented strains were spotted onto plates containing YNB and oleate or glucose (as a control) as the carbon source. All of the strains grew well on glucose; however, the cat2Δ/cat2Δ strain was unable to grow on oleate (Fig. 2), a phenotype that was confirmed by carrying out growth assays in liquid media (data not shown). These results support the hypothesis that C. albicans is dependent on the activity of Cat2p to export acetyl units produced during fatty acid β-oxidation and confirm that C. albicans lacks a peroxisomal citrate synthase. As the cat2Δ/fox2Δ strain lacks both the peroxisomal and mitochondrial forms of Cat2p, it is unclear what each form contributes to the phenotype on oleate. Remarkably, the cat2Δ/fox2Δ strain is also not able to grow on the C2 carbon source acetate or ethanol (Fig. 2), a condition under which acetyl-CoA is produced in the cytosol. Together, these results show that both the peroxisomal and mitochondrial forms of Cat2p are essential for growth on nonfermentable carbon sources.

The cat2 null mutant shows reduced β-oxidation activity and elevated levels of acetyl-CoA. To study the oleic acid metabolism of the cat2 null mutant in more detail, we determined its total β-oxidation activity by incubating intact cells with 1-14C-labeled oleic acid and measuring the labeled CO2 and acid-soluble counts produced (representing carbon metabolism intermediates) (39). The C. albicans fox2Δ/fox2Δ strain, lacking the second enzyme of the β-oxidation pathway, served as a negative control, as this mutant has virtually no fatty acid β-oxidation activity (26). The fox2Δ/fox2Δ and cat2Δ/cat2Δ strains showed greatly reduced production of CO2 compared to the wild-type and CAT2 complemented strains (Fig. 3A). The incorporation of radiolabel into acid-soluble material was very low in the fox2Δ/fox2Δ strain, but significant amounts were found in the cat2Δ/cat2Δ strain, suggesting that the labeled fatty acid can still be converted to acetyl-CoA and/or other carbon metabolism intermediates (39). The C. albicans fox2Δ/fox2Δ strain, lacking the second enzyme of the β-oxidation pathway, served as a negative control, as this mutant has virtually no fatty acid β-oxidation activity (26). The fox2Δ/fox2Δ and cat2Δ/cat2Δ strains showed greatly reduced production of CO2 compared to the wild-type and CAT2 complemented strains (Fig. 3A). The accumulation of acetyl-CoA in the cat2 mutant may account for the relatively large amount of acid-soluble material produced in this strain compared to the fox2
null strain (Fig. 3B). Together, these data show that acetyl-CoA transport from peroxisomes to mitochondria is defective in cells lacking Cat2p.

The cat2 null mutant is not attenuated in virulence in the mouse model of systemic candidiasis. Previously, it was shown that carnitine acetyltransferase plays an essential role during plant infection by the rice blast fungus *M. grisea* (2). We used the mouse model of systemic infection to test the role of Cat2p in the virulence of *C. albicans*. BALB/c mice were injected intravenously with cells of the wild-type, cat2Δ, or complemented strain at a challenge dose of 1.5 × 10⁴ CFU/g body weight. No significant differences were found between the strains with respect to survival times (as calculated by log rank test) or tissue burdens (t test) (Fig. 4A and B). These results indicate that carnitine acetyltransferase does not play an essential role in *C. albicans* infection in the mouse model.

The cat2 null mutant shows a small but significant defect in biofilm formation. We tested whether the absence of the metabolic enzyme Cat2p affected biofilm formation over a period of 72 h and observed that significantly less biofilm was formed by the cat2 null mutant compared to the wild-type and complemented strains as quantified by crystal violet staining (Fig. 5A). Direct measurement of biofilm thickness with an LLCT (25) revealed that the cat2 null mutant formed 30% thinner biofilms compared to the wild-type strain (350 and 250 μm thick, respectively) and that biofilm formation was partially restored in the complemented strain (300 μm thick), which is in line with the gene dosage effect (Fig. 5B). CLSM analysis revealed no difference among the three strains with respect to biofilm architecture or morphology and confirmed the reduced ability of the cat2 null mutant to form biofilm (Fig. 5C).

**DISCUSSION**

Two possible pathways for the transport of acetyl units from peroxisomes to mitochondria have been described in plants and yeasts, of which one is reliant on peroxisomal citrate synthase and the other is dependent on peroxisomal carnitine acetyltransferase. Here we show that the human fungal pathogen *C. albicans* lacks a peroxisomal citrate synthase and exclusively relies on Cat activity for the export of acetyl units from peroxisomes. We previously showed that the Cit and Cat routes work in parallel since deletion of either peroxisomal citrate synthase (*cit2Δ*) or the major Cat protein (*cat2Δ*) does not affect growth on oleate but deletion of both does (39). Current evidence suggests that plant peroxisomes lack carnitine acetyltransferase and use the
citrate synthase pathway to shuttle acetyl units to mitochondria (27). Thus, whereas S. cerevisiae can employ both pathways, A. thaliana and C. albicans are each dependent on a single pathway, Cit in A. thaliana and Cat in C. albicans (Fig. 6). A BLAST search of the sequenced fungal genomes (http://www.broad.mit.edu/annotation/fungi and http://cbi.labri.fr/Genolevures/BLAST.php) revealed that at least three other closely related fungal species (Candida tropicalis, Debaryomyces hansenii, and Candida lusitaniae) contain a single CIT gene predicted to encode a mitochondrion-targeted citrate synthase, suggesting that they, like C. albicans, are dependent on peroxisomal Cat activity for growth on fatty acids. The fact that C. albicans grows very well on mineral oleate medium lacking carnitine (Fig. 2 and our unpublished observations), together with the observed strict requirement for Cat activity under these growth conditions, implies that C. albicans possesses a functional carnitine biosynthesis pathway. Experiments to provide evidence of this are under way in our laboratory.

We have shown that the cat2Δ/cat2Δ strain not only is unable to grow on fatty acids but also does not grow on ethanol or acetate (Fig. 2). Under the latter conditions, the acetyl-CoA is produced in the cytosol. Since the peroxisomal and mitochondrial membranes are impermeable to acetyl-CoA, it is conceivable that the acetyl units must be linked to carnitine to allow their transport over the peroxisomal and mitochondrial membranes and enter the glyoxylate cycle and TCA cycle, respectively. The absence of the mitochondrial and peroxisomal Cat enzymes in the cat2 null mutant prevents the reformation of acetyl-CoA inside the organelles and thus further metabolism. Which enzyme(s) is involved in linking the acetyl units to carnitine in the cytosol is unclear. This function may be carried out by either of the two other predicted CAT genes, CTN1 and CTN3, which are potential homologs of S. cerevisiae YAT1 and YAT2, respectively (28). In S. cerevisiae, both YAT genes are expressed on ethanol and Yat1p was shown to be associated with the outer mitochondrial membrane, suggesting that Yat1p and/or Yat2p may be involved in the generation of acetyl-carnitine in the cytosol (33, 35). The subcellular localization of Ctn1p and Ctn3p in C. albicans has not been addressed experimentally, but the absence of a clear mitochondrial or peroxisomal targeting signal 1 (http://ihg.gsf.de/ihg/mitoprot.html and http://mendel.imp.ac.at/mendeljsp/sat/pts1/PTSPredictor.jsp) suggests that they may function in the cytosol. However, no significant amounts of Cat activity could be detected in the cat2 null mutant grown on glucose, acetate, ethanol, or oleate (Table 4), despite our efforts to measure this activity under a variety of assay conditions. It remains possible, however, that Ctn1p and Ctn3p are not active in our assay or that they are expressed under very specific conditions. Indeed, Prigneau et al. (28) have shown by Northern blot analysis that CTN1, CTN3, and CAT2 transcripts are induced during macrophage infection but enzyme activities were not determined. How the cytosolic acetyl-CoA produced enters the peroxisomal and mitochondrial compartments remains therefore unresolved.

Previously, we and others investigated the role of fatty acid metabolism in the virulence of C. albicans (19, 26). Piekarska et al. showed that a C. albicans fox2Δ/fox2Δ strain lacking the second enzyme of the β-oxidation pathway shows attenuated virulence in mice but that the virulence defect of this strain is probably caused by a dysfunctional glyoxylate cycle (26). The phenotype of the cat2 null mutant is similar to that of the fox2Δ/fox2Δ strain in that it exhibits greatly reduced β-oxidation activity and an inability to grow on oleate, acetate, and ethanol. However, the glyoxylate cycle in the oleate-grown cat2Δ/cat2Δ strain may still be partially functional, as can be inferred from the conversion of fatty acids into acid-soluble material representing carbon metabolism intermediates (Fig. 3).

The wild-type virulence of the cat2 strain in the mouse model corroborates previous data obtained with the fox2Δ/fox2Δ strain and shows independently that fatty acid β-oxidation is not required for the survival of C. albicans in infected mice (Fig. 4). In line with this, Barelle et al. recently reported that the dominant metabolic mode of C. albicans single cells in vivo is glycolytic rather than gluconeogenic (1).

Carnitine acetyltransferase does play an essential role in the virulence of the plant-pathogenic fungus M. grisea, as shown by the inability of a pth2 mutant lacking the major Cat enzyme to form penetration hyphae and cause plant infection (2, 30). Interestingly, the pth2 mutant is also not able to grow on fatty acids (olive oil) or acetate, indicating that M. grisea, like C. albicans, is dependent on the Cat pathway for growth on these carbon sources. By using green fluorescent protein (GFP)-tagged versions of Pth2p, a unique peroxisomal localization was found for the protein (2). Since the Pth2p ortholog Cat2p in S. cerevisiae and C. albicans has dual localizations and contains targeting signals in both the N-terminal (mitochondrial) and C-terminal (peroxisomal) parts of the protein, it is very well possible that GFP tagging of Pth2 has disturbed its mitochondrial targeting, causing it to localize only to peroxisomes. However, the Pth2-GFP construct did fully complement the pth2 mutant phenotype, suggesting that peroxisome-targeted Pth2p is functional. Determination of the subcellular distribution of the endogenous untagged protein is required to resolve this issue.

Cell-cell adherence, the ability to form hyphae, and production of an extracellular matrix are required for robust biofilm formation by C. albicans (3, 5). Because a C. albicans ctn3 null mutant showed reduced hypha formation (28) and an M. grisea pth2 mutant showed an increased sensitivity to cell wall stress (2, 30), we determined whether our cat2 null mutant exhibited similar phenotypes. No noticeable differences in sensitivity to cell wall-perturbing compounds (Calcofluor white, Congo red, sorbitol, and sodium dodecyl sulfate) or the ability to form hyphae were found between the wild type and the null mutant (data not shown). However, a small but significant reduction in the ability to form biofilms in vitro was observed in the cat2 deletion mutant (Fig. 5), possibly because the mutant is unable to metabolize nonfermentable carbon sources, which may cause a growth disadvantage in the later, mature stages of biofilm formation when glucose is exhausted. Further experiments are required to address this issue.

In conclusion, our studies revealed that C. albicans exclusively relies on Cat activity for the transport of acetyl units among peroxisomes, cytosol, and mitochondrion during growth on nonfermentable carbon sources. While the lack of Cat activity does not affect the virulence of C. albicans in mice, maximum biofilm formation does require a functional Cat pathway. Following the observations of Mukherjee and colleagues on the role of alcohol dehydrogenase (22), Cat2p is
now the second example of a metabolic enzyme affecting biofilm formation in \textit{C. albicans}.

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**ADDENDUM**

While the manuscript was under review, a paper reporting similar data was published by the group of Lorenz (40).

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