A *Histoplasma capsulatum* lipid metabolic map identifies antifungal targets and virulence factors

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Running title: *H. capsulatum* lipid metabolism

Keywords: *Histoplasma capsulatum*, lipid metabolism, drug target, platelet-activating factor, multi-omics analysis
Abstract

Lipids play a fundamental role in fungal cell biology, being components of the cell membrane as well as targets of antifungal drugs. A deeper knowledge of lipid metabolism is key for developing new drugs and a better understanding of fungal pathogenesis. Here we built a comprehensive map of the *Histoplasma capsulatum* lipid metabolic pathway by integrating proteomic and lipidomic analyses. The map led to the identification of both the fatty acid desaturation and the sphingolipid biosynthesis pathways as targets for drug development. We also found that *H. capsulatum* produces analogs of platelet-activating factor, a potent regulator of the human immune response. The *H. capsulatum* platelet-activating factor analogs induced platelet aggregation and stimulated the production of the cytokines interleukin-10 and tumor necrosis factor-α by J774 macrophages. Overall, this lipid metabolic map revealed pathways that can be targeted for drug development, in addition to identifying a regulator of the host immune response.
Introduction

Fungal diseases affect more than 1 billion people and cause 1.6 million deaths every year (1). *Histoplasma capsulatum*, the causative agent of histoplasmosis, is an important pathogen in this context, being associated with HIV infections and a major cause of morbidity and mortality worldwide (2). A 2018 CDC report, reviewed hospitalization data from 2011-2014 in only 12 states in the US and identified 3,409 cases of histoplasmosis with a 7% mortality rate (3). Serological surveys have shown that sera from 60-90% of individuals are reactive to the *H. capsulatum* antigen histoplasmin in communities surrounding the Mississippi and Ohio basins, suggesting that epidemiological numbers of the disease could be underestimated (4, 5).

Histoplasmosis treatment relies on only a few antifungal drugs, but the increasing number of resistant strains is a major public health concern (6, 7). The absence of immunotherapies and the fact that the newest class of antifungal drugs, echinocandins, has already been available for two decades, make the development of new therapies a high priority. A major hurdle in developing new drugs is the limited knowledge about the detailed metabolic reactions and mechanisms of pathogenesis of this microbe.

Lipids have essential roles in many biological processes and the biosynthetic pathways of fungal lipids diverged from metazoan pathways, which makes them obvious antifungal drug targets (8). Indeed, two classes of current antifungal drugs target lipid biosynthesis: 1) polyenes, including amphotericin B and nystatin, bind to ergosterol causing increased fungal cell membrane permeability and loss of intracellular constituents; and 2) azoles, such as voriconazole, itraconazole and fluconazole, inhibit the cytochrome P-450-dependent 14α-sterol demethylase, which is an enzyme of the ergosterol biosynthetic pathway (8, 9). More recently, glucosylceramide, a type of sphingolipid that is critical for infection of many fungal species, has been validated as a drug target in *Cryptococcus neoformans* (10, 11). Lipids have also been characterized as important molecules for fungal infections (12). For instance, glycolipids, sterols, as well as the chain length and level of desaturation of fatty acids play a role in pathogenesis (11, 13-18). Not only the specific lipid species but how they are organized into larger structures are also important for virulence. For instance, ergosterol-dependent microdomains have a crucial role in infectivity (19, 20). The role of lipids in virulence opens an opportunity to design drugs that target pathogenesis processes rather than the viability of fungal cells (21).

There is limited information about *H. capsulatum* lipid composition and function. The levels of *H. capsulatum* phosphatidylcholine correlate with its virulence (22). Similarly, perturbations in *H. capsulatum* membrane lipids reduce virulence (14, 23) and alterations in a
delta-9 fatty acid desaturase gene enhance temperature sensitivity (24). Also, unsaturated fatty acids decrease in aged yeast cells (25). In terms of sphingolipids, *H. capsulatum* also produces glucosylceramides, inositolphosphorylceramides and galactosyl-dimannosyl-inositolphosphorylceramides, which are components of membrane microdomains and are important for infection (20, 26, 27). Considering the importance of lipids, we reasoned that a deeper characterization of *H. capsulatum* lipid metabolism would result in the discovery of virulence factors and drug targets. In this study, we performed an in-depth characterization of the *H. capsulatum* lipid biosynthetic pathway by profiling its lipids and the associated proteins, which were integrated into a metabolic map. By comparing our results to *Saccharomyces cerevisiae* and humans, we show unique features of the lipid biosynthetic pathway of *H. capsulatum* that can be targeted for drug development. This analysis also led to the discovery that *H. capsulatum* produces analogs of platelet-activating factor (PAF), which are biologically active and regulate different aspects of the human immune response.
Results

Proteomic and lipidomic analyses and overview of the *H. capsulatum* lipid metabolism

To determine the global landscape of the lipid metabolic pathway of *H. capsulatum* we performed comprehensive lipidomic and proteomic analyses. Samples were extracted using two different methods and submitted to chromatographic separation before being analyzed by mass spectrometry. Paired global lipidomics and proteomics analyses were performed by submitting the samples to simultaneous metabolite, protein and lipid extraction (MPLEx)(28) (Fig. 1a). For complementary analyses of sterols, fatty acids and phospholipids, yeast cells were extracted with two rounds of organic solvent extraction, followed by a solid phase fractionation in a silica 60 column (Fig. 1a). The extracted fractions were analyzed by either gas chromatography-mass spectrometry (GC-MS) (sterols and fatty acids) or liquid chromatography-tandem mass spectrometry (LC-MS/MS) (sphingolipids, glycerolipids, phospholipids and proteins) (Fig. 1a).

The combined analysis identified 371 unique lipid species from 5 major lipid categories (fatty acids, sterols, glycerolipids, sphingolipids and glycerophospholipids (phospholipids + lysophospholipids)) that were subdivided into 19 subclasses (Fig. 1b and Supplementary Tab. 1-8). The most diverse subclasses of lipids in terms of number of identified lipids were triacylglycerols (TG), phosphatidylcholines (PC) and phosphatidylethanolamine (PE) with 130, 57 and 51 individual species, respectively (Fig. 1b). The proteomic analysis led to the identification of 3,215 proteins (Supplementary Tab. 9). To provide a measurement of the protein abundances in the cells, we calculated the relative copy number of each protein and scaled them into high, moderate, low and very low abundance (Fig. 1c), using a scale similar to one previously described (29). To validate this scale, we performed a function-enrichment analysis using the KEGG annotation to check the abundance of different pathways. As expected, glycolysis/gluconeogenesis and tricarboxylic acid (TCA) cycle were overrepresented among the highly abundant proteins, whereas DNA replication, nicotinate and nicotinamide metabolism, and basal transcription factors were enriched in moderate, low and very low abundance levels, respectively (Fig. 1d). The same scale showed that fatty acid metabolism was enriched in highly abundant proteins, whereas steroid biosynthesis proteins were mainly present in moderate abundance (Fig. 1c-d). Glycerophospholipid and sphingolipid metabolism proteins were not concentrated in a single abundance level and were spread mostly between moderate and low abundant levels (Fig. 1c-d). The abundance levels of the proteins were directly proportional to each type of lipid in the cellular membrane. For instance, the proteins of fatty acid and steroid (the building blocks of the cell membrane) metabolism were more abundant compared to proteins of the glycerophospholipid, ether lipid or sphingolipid
metabolism, which are responsible for the synthesis lipids from specific lower-abundance classes (Fig. 1d).

**Lipid biosynthesis and remodeling map of *H. capsulatum***

To provide a global view of *H. capsulatum* lipid metabolism we built a map including lipid biosynthesis and remodeling reactions. The map was constructed based on a previous publication (30) and conserved metabolism between *H. capsulatum* vs. *S. cerevisiae* and *Cryptococcus neoformans*, the best characterized fungal organisms in terms of lipid metabolism (Supplementary Tab. 10). The *H. capsulatum* map was integrated with the relative abundance of the lipid species within the same subclass and the protein abundances (Fig. 2). In terms of fatty acids, lipids containing 16 and 18 carbons were the most abundant. Consistent with figure 1D, 5 out of 10 proteins of this pathway were highly abundant (Fig. 2). Similarly, 13 of the 23 sterol biosynthesis proteins had moderate abundance, with ergosterol being the most abundant product (Fig. 2). In the sphingolipid pathway, ceramides (Cer), hexosylceramide (HexCer), and inositolphosphoceramides (Pl_Cer) were the detected lipid species (Fig. 2). Out of the 9 proteins detected in the proteome, 3 had low abundance, 5 had moderate abundance and 1 was highly abundant (Fig. 2). The number of low abundant proteins does not necessary mean that the pathway has low activity. For instance, one low abundant protein had more abundant paralogues with the same function (Lcb2 vs. Lcb1), and the other two proteins regulate the specific modification of the head group of HexCer (Fig. 2). In terms of glycerolipids, diacylglycerols (DGs) and TGs were identified, being the proteins from this pathway that were present at moderate (2), low (3) and very low abundance (1) (Fig. 2). Like the free-fatty acid composition, the most abundant species of DGs and TGs had fatty acyl groups with either 16 or 18 carbons attached to them (Fig. 2). Consistent to the glycerolipids, all the different glycerophospholipid classes had species bearing C16 and C18 as the most abundant in each (Fig. 2).

We further curated the map by studying the substrates and products of the acyltransferase Ale1 (also known as lysophosphatidylcholine (LPC) acyltransferase 1 – LPT1). A lipidomic analysis was performed in the *Lpt1* knockout strain of *S. cerevisiae* complemented or not with the *H. capsulatum* (*Hc-Lpt1*) orthologue (53% similarity to *S. cerevisiae* Lpt1 – *Sc-Lpt1*), using a plasmid with a galactose-inducible promoter. Both gene knockout and recombinant expression were validated by proteomic analysis (Fig. 3a). As expected, the *Sc-LPT1* specific peptide DISASSPNLGGILK was detected in wild-type *S. cerevisiae*, in both glucose- and galactose-supplemented media (Fig. 3a). The *Hc-LPT1*-specific peptide
LTAFCWNVHDGR was detected only in complemented strains that were grown in galactose-supplemented medium (Fig. 3a). The analysis showed that Hc-LPT1 complementation increased the abundances of 60 lipids, including phosphatidic acid (PA), PC, PE, phosphatidylinositol (PI), PS, DG, TG, Cer and mannosyl-inositolphosphoceramide (MIPC) (Fig. 3b-d, Supplementary Table 11). The Hc-LPT1 complementation also significantly reduced the levels of 77 lipids, including 5 LPC, 3 lysophosphatidylethanolamines (LPE), 2 sphinganines and 3 diacylglycerols (Supplementary Tab. 12). On the other hand, disruption of the Sc-LPT1 gene reduced the levels of 30 potential products, including PC, PE, PI, PS and TG. Notably, the proportion of acylated products between the LPT1 from the two species was very different, being PC, PE, PI and PS being more acylated by the S. cerevisiae ortholog and PA, DG, TG, Cer and MIPC by the H. capsulatum counterpart (Fig. 3D, Supplementary Tab. 13). The Sc-LPT1 showed a preference to produce lipid species containing short fatty acyl chains, with all the identified products having chains with ≤ 14 carbons (Fig. 3c). Complementation with Hc-Lpt1 gene showed a different phenotype, as this orthologue produced lipid species containing odd-carbon number (C15 and C17) and longer (> C18) fatty acyl chains (Fig. 3e). Overall, the map provides a global view of lipids being produced by H. capsulatum along with catalytic enzymes.

**Divergent lipid metabolic pathways as drug targets**

We next focused on pathways that could be targeted for antifungal chemotherapies. We searched for divergent pathways comparing H. capsulatum with S. cerevisiae and humans as candidates for anti-H. capsulatum drug targets. The most divergent pathways were fatty acid and sphingolipid metabolism. We examined the distribution of fatty acyl chains that are incorporated into H. capsulatum and S. cerevisiae. The results showed that both fungi incorporate fatty acyl groups with different chain lengths into their lipids. S. cerevisiae incorporate significantly more fatty acyl chains with less than 16 carbons (by Fisher’s exact test), whereas H. capsulatum has significantly more lipids with fatty acids longer than 17 carbons (Fig. 4a). In terms of unsaturation, S. cerevisiae had almost exclusively saturated lipids or fatty acyl chains with 1 double bound, whereas H. capsulatum also had fatty acyl chains with 2 or 3 double bonds (Fig. 4b-c). This is consistent to the fact that S. cerevisiae has only one fatty acid desaturase gene (delta-9 desaturase, also known as oleate synthase Ole1) in its genome. In addition to Ole1, H. capsulatum has a delta-12 desaturase (FAD2) and two uncharacterized desaturases (HCBG_08514 and HCBG_05092) that might be responsible for the conversion of C18:2 and C20:2 to C18:3 and C20:3, respectively (top left part of Fig. 2). Because the human genome lacks delta-9 and delta-12 desaturases (Fig. 4b), we decided to
test it as a potential drug target. We tested two fatty acid desaturase inhibitors, 10-thiastearic acid and thiocarlide, as growth inhibitors of *H. capsulatum* yeasts. 10-Thiastearic acid showed a minimum-inhibitory concentration (MIC) of 1.25 μM, whereas the thiocarlide MIC was 12.5 μM (Fig. 4d). To confirm that these compounds target fatty acid desaturases we performed a fatty acid analysis. Cells were grown for 48 hours with one half of the MIC and fatty acids were extracted, methylated and analyzed by GC-MS. 10-Thiastearic acid had no effect on the ratio between unsaturated and saturated fatty acids (Fig. 4e), suggesting that this compound might target other pathways rather than the fatty acid desaturation. As expected, thiocarlide reduced the C18:2/C18:0 peak area ratio from 11.4 to 8.5 (25% reduction), the C18:1/C18:0 ratio from 4.5 to 3.7 (18% reduction) and the C20:1/C20:0 ratio from 1.3 to 0.8 (39% reduction) (Fig. 4e). Conversely, the C16:1/C16:0 ratio had a slight increase from 0.036 to 0.044 (22% increase) (Fig. 4e). These results show that the unsaturated fatty acid pathway can be targeted for developing anti-histoplasmosis drugs.

Sphingolipid metabolism was another pathway that had major differences in *H. capsulatum* compared to *S. cerevisiae*. *H. capsulatum* produces hexosylceramides (glucopyranosylceramides)(27), which are absent in *S. cerevisiae*. Even though both fungi produce inositolphosphoceramides and mannosyl-inositolphosphoceramides, they branch into different products (Fig. 5a). Whereas *S. cerevisiae* mannosyl-diinositolphospho-ceramides as a final product, *H. capsulatum* produces galactosyl-dimannosyl-inositolphosphoceramides (31) (Fig. 5a). Also, *H. capsulatum* makes two major pools of ceramide (4-hydroxysphinganine and 9-methyl-4,8-sphingadienine) but *S. cerevisiae* only makes 4-hydroxysphinganine (27, 30). Compared to fungi, humans produce a very distinct profile of sphingolipids. Unlike end product monohexosylceramides of fungi, humans synthesize a much more complex set of glycosphingolipids, including globo, isoglobo, lacto and gangliosides (32) (Fig. 5a). Conversely, fungi produce inositolphosphoceramides and their glycosylated derivates, which are absent in humans. Humans also produce sphingomyelins, which are absent fungi (Fig. 5a). The sphingolipid pathway can therefore be an excellent drug target due to these major differences in composition compared to humans. We performed a proof-of-concept by inhibiting the first enzyme of the sphingolipid biosynthesis, serine-palmitoyltransferase, with myriocin. Myriocin had a MIC of 30 nM (Fig. 5b). To confirm that myriocin targets the sphingolipid pathway, lipidomic analysis was performed on cells grown with one half of the MIC for 48 hours. As expected, the most abundant ceramide species, Cer(d18:1/24:0(2OH)) and Cer(d20:0/24:0), were reduced by 52% and 62%, respectively (Fig. 5c). As a control of a lipid from an unrelated class, the level of the most abundant DG, DG(16:0/18:2/0:0), increased by 51% (Fig. 5c). These
results validate the sphingolipid pathway as a potential target for developing anti-\textit{H. capsulatum} drugs.

**Discovery and validation of platelet-activating factor analogs in \textit{H. capsulatum}**

One unique lipid found in \textit{H. capsulatum} was PC(16:2/2:0) (Fig. 6a and Supplementary Fig. 1). This lipid is analogous to the canonical platelet activating factor (PAF), a PC species containing ether-linked C16:0 chain in the sn1 position and acetyl group in the sn2 position (2-O-acetyl-1-O-hexadecyl-sn-glycero-3-phosphocholine – PC(O-16:0/2:0)) (Fig. 6a). PAF is a major regulator of the immune response in mammals (33). Therefore, we sought to further validate the structure of the \textit{H. capsulatum} PAF analog. We searched for other lipids containing acetate as a fatty acyl chain in our lipidomics data and found another species, PC(16:1/2:0) (Supplementary Fig. 2). We next aimed to compare our molecule to a chemically synthesized standard, but unfortunately, PC(16:2/2:0) and PC(16:1/2:0) standards were not commercially available. The lack of knowledge on the double bond position prevented to synthesize the lipid containing the right fatty acyl group. We reasoned that \textit{H. capsulatum} could be producing a saturated version of these lipid species (1-palmitoyl-2-acetyl-sn-glycerol-phosphocholine – PC(16:0/2:0) which is commercially available) (Fig. 6a), although in smaller amounts. The liquid chromatography-tandem mass spectrometry analysis found a peak with the expected \textit{m/z} (mass to charge ratio) and retention time (Fig. 6b). To ensure that the sample background would not cause retention time shift, the sample was spiked with the standard and analyzed in parallel, which showed a consistent elution pattern with the standard alone and the endogenous species (Fig. 6b). Tandem mass fragmentation of this species showed a major fragment at \textit{m/z} 184.07, derived from the breakage of the phosphocholine (ChoP) head group (Fig. 6c). Because tandem mass fragmentation of lipid species often generates only a few fragments, which is not optimum to confidently identify a molecule, we performed ion-mobility spectrometry coupled with mass spectrometry (IMS-MS) analysis. In the drift tube IMS-MS analyses performed, the ions were gently pulled by a weak electric field through the drift tube that is filled with nitrogen gas. As the ions collided with the gas molecules, they were separated based on their shape and size, allowing the separation of isobaric molecules which cannot be distinguished by \textit{m/z} alone. \textit{H. capsulatum} lipid extract was fractionated using a C18 column coupled to a HPLC system and fractions enriched with the lipids of interest were collected and analyzed by IMS-MS. The results showed a consistent drift time and collision cross section values when comparing the endogenous PC(16:0/2:0) to the synthetic standard (Fig. 6d). Overall, the retention time, high
mass accuracy, tandem mass fragmentation and drift time of the IMS-MS analyses consistently support the presence of acetylated PCs in *H. capsulatum*.

**H. capsulatum PAF analogs are biologically active**

To determine if the *H. capsulatum* PAF analogs have similar activity compared to the human canonical PAF, yeast lipid extracts were submitted to solid-phase extraction in a Silica 60 column and further separated by HPLC reverse phase chromatography using a C18 column. Each fraction was monitored by IMS-MS and the fractions containing *H. capsulatum* PAF analogs (fractions 8-10 in Fig. 7) were tested for their ability to aggregate platelets. Platelet aggregation was measured by impedance in whole blood. The results showed that the *H. capsulatum* fractions containing PAF analogs were active and promote the platelet aggregation, like the positive controls with the canonical PAF, PC(16:0/2:0) (acyl-PAF) and arachidonic acid (Fig. 7a-b).

We also tested the HPLC fractions enriched in *H. capsulatum* PAF analogs in their ability to induce cytokine production in J774 macrophage-like cells. The fractions induced a dose-dependent production of interleukin-10 (IL-10) and tumor necrosis factor (TNF)-α in pico to nanomolar range (estimated based on peak area relative to external standards), which is consistent to the activity of the PC(16:0/2:0) standard (Fig. 8a-b). The highest tested concentration of the lipid fractions and standard induced the production of a smaller amount of TNF-α (Fig. 8a-b), probably due to saturation or negative feedback mechanisms. The TNF-α production was inhibited by the PAF receptor antagonist WEB2086 (Fig. 8a-b), showing that *H. capsulatum* PAF analogs regulate J774 cells by engaging the PAF receptor.

Taken together our data show that *H. capsulatum* produces bioactive PAF analogs that activate platelets and induce cytokine production in host effector cells.
Discussion

Here we developed a global map of the *H. capsulatum* lipid metabolism by integrating genomic, proteomic and lipidomic information, including relative abundances of proteins and lipids. We found an accordance of the protein abundance levels with their function in the lipid metabolism. For instance, the proteins involved in the synthesis of the major lipid components, the fatty acids and sterols, were enriched among the high and moderate abundant proteins, whereas the proteins related to the synthesis of specific lipid head groups were enriched in the low to very low abundant proteins. This might be a consequence of cell resource optimization as protein synthesis is one of the most energetically expensive tasks in cells (34). Our lipid map also showed that TG, PC and PE are the most diverse classes of lipids. This could reflect the abundance of these lipid classes, as they are the most abundant ones(35). Our results show that the diversity of TG, PC and PE species could be due to fatty acid remodeling, as these lipids are products of the LPT1 acyltransferase. LPT1 transfers a variety of fatty acyl chains, including odd-carbon chain (C15 and C17) and long chain (≥C18) fatty acids. Our experiments with LPT1 also helped to curate the lipid map since the *S. cerevisiae* orthologue has a different specificity for substrates, that being a major impact in producing phospholipids with short fatty acyl chains. In *S. cerevisiae*, LPT1 has been shown in vitro, using radioactive precursors to acylated LPC, LPG, LPA, LPE, LPI and LPS (36). The acylation of LPA, a central precursor for all glycerolipids and glycerophospholipids, suggest that the LPT1 impact could be indirect. Therefore, further investigation will be needed to determine if this enzyme can directly acylate DG into TG.

We identified the fatty acid desaturation and the sphingolipid pathways as divergent points in the lipid metabolism pathways of *H. capsulatum* vs. *S. cerevisiae* and humans. We showed that these two pathways are potential targets for developing anti-histoplasmosis drugs. Thiocarlide (also known as isoxyl) is a potent inhibitor (nanomolar range) of *Mycobacterium tuberculosis* delta-9 stearoyl desaturase and was used as a second line of anti-tuberculosis drugs in the 1960s (37, 38). However, despite its low toxicity, thiocarlide was not as effective during clinical trials, so is no longer used to treat patients (38-40). One possible explanation for the clinical trial failure is the poor solubility of the compound in water. Since then, other drug delivery vehicles have been tested with promising results in vitro (41). Delta-12 oleate desaturase has been validated as a drug target in trypanosomes (42). Indeed, both 10-thiastearic acid and thiocarlide inhibit *Trypanosoma cruzi* growth in low micromolar range (43), similar to the MICs for *H. capsulatum* (Fig. 4d). Although toxicity has not been tested directly with 10-thiastearate, its analog tetradecylthioacetic acid has a low toxicity in humans up to
1g/day (44), reinforcing the potential of this class of molecules as antimicrobial drugs. Our data showed that 10-thiastearate might target a different pathway rather than fatty acid desaturation, but it should still be considered a drug candidate due to its antifungal activity and possible low toxicity to humans.

We also performed a proof-of-concept that sphingolipids are promising targets for anti-histoplasma drug development by inhibiting the first step of the sphingolipid pathway with myriocin. Myriocin kills *Candida albicans* and *Aspergillus fumigatus* in micromolar concentrations (45, 46). *H. capsulatum* is more sensitive to myriocin as its MIC was 30 nM (Fig. 5b). Unfortunately, myriocin has immunosuppressant activity (47, 48), but this has been explored for simultaneously killing the fungus and reducing the pathogenic inflammation of the lungs in cystic fibrosis mice infected with *A. fumigatus* (49). The fact the de novo sphingolipid synthesis is essential for *H. capsulatum* opens the opportunity to explore other inhibitors of serine palmitoyltransferase, or other enzymes in the pathway. Indeed, other enzymes of the fungal sphingolipid pathway have been validated as a drug target. In *C. neoformans*, acylhydrazones have been shown to be potent inhibitors of the glucosylceramide synthesis (nanomolar range) and excellent antifungal drug candidates (10, 50).

Our study also led to the identification of PAF analogs in *H. capsulatum*. Such analogs were previously described in *S. cerevisiae*, which is synthesized by the acyltransferase LTP1 (36, 51). We were able to validate these findings by analyzing the Sc-LPT1 knockout strain, which had the production of PC(16:1/2:0) drastically reduced (Supplementary Fig. 3). Unfortunately, the *H. capsulatum* preferred substrate LPC(16:2) is not present in *S. cerevisiae*, due to the lack of additional desaturases in this species. Complementation of *S. cerevisiae* knockout strain with the Hc-LPT1 orthologue significantly increased the abundance of PC(16:1/2:0) (Supplementary Fig. 3). Therefore, it is possible that LPT1 is PAF synthase in *H. capsulatum*. However, this and other possibilities, such as PAF being synthesized by a different acyltransferase, need to be confirmed with more experiments. The *H. capsulatum* PAF analogs induced platelet activation and PAFR-dependent cytokine production. PAF has a variety of functions in regulating the host immune response and it has been shown to be essential for neutrophil recruitment to the site of infection in histoplasmosis (52). Since acyl-PAF has pro- and anti-PAF activity depending on the physiological condition (53), it is possible that the *H. capsulatum* PAF analogs play a role in controlling the immune response against histoplasmosis.

In conclusion, here we built a detailed map of *H. capsulatum* lipid metabolism. Our data identified regions of the *H. capsulatum* lipid metabolic pathway that can be targeted for drug
development and led to the characterization of lipid species that have immunoregulatory activities.
MATERIAL AND METHODS

Fungal cell culture

*Histoplasma capsulatum* G217B strain was purchased from the American Type and Culture Collection (ATCC, Manassas, VA) and grown in yeast form at 37°C in Ham’s F12 medium as previously described (54). *S. cerevisiae* (MATalpha background) deficient LPT1 gene and wild-type strains were purchased from Dharmacon (Lafayette, CO). The strains were transformed by a lithium-acetate-heat shock method with a plasmid containing the Hc-LPT1 gene (Uniprot ID C0NZS2) that was codon optimized, synthesized and clone into the Gal-inducible plasmid pESC-URA by GenScript (Piscataway, NJ). The cells were cultivated in YNB medium containing leucine, lysine, histidine and uracil, further supplemented with of glucose or galactose at 28 °C with shaking at 200 rpm until the culture reached an OD600 of 0.8-1.0. Cells were harvested by centrifuging at 2000 x g for 2 min and washed twice with PBS. All the sample preparation and analysis were done in 4 replicates and in a randomized order to ensure the biological significance and prevent batch effects.

Sample extraction and fractionation on Silica 60 column

Samples were extracted using two different approaches. For global proteomics and lipidomics analyses, samples were submitted to Metabolite, Protein and Lipid Extraction (MPLEx) as previously described (28). For the fractionation study, cells were suspended in 400 μL of water and transferred to PTFE-lined 13x100-mm Pyrex glass tubes (prewashed with chloroform:methanol (2:1, v:v) and chloroform:methanol:water (1:2:0.8, v:v:v)) containing 200 μL of silica/zirconia beads (55). 1 mL of methanol and 0.5 mL of chloroform to form chloroform:methanol:water (1:2:0.8, v:v:v) solution and extracted by vortexing for 2 minutes followed by centrifugation for 10 minutes at 1,800 x g at room temperature. The supernatants were collected into clean glass tubes. Samples were additionally extracted twice with 1 mL of chloroform:methanol (2:1, v:v) and once with 1 mL of chloroform:methanol:water (1:2:0.8, v:v:v), and the supernatants were pooled together then dried in a vacuum centrifuge. Extracted lipids were fractionated into neutral lipids, fatty acids and phospholipid fractions using Silica 60 columns (56) and dried in a vacuum centrifuge.

High performance liquid chromatography (HPLC) fractionation of phospholipids

The phospholipid fraction from the Silica 60 separation was reconstituted in 2 mL methanol:dichloromethane (8:2), centrifuged and the supernatant was loaded (100 μL) onto a
reverse-phase column (Kinetex®, Torrance, CA, 150 x 4.6 mm, 2.6 µm, C18, 100 Å,) coupled to a 1200 Agilent HPLC and fraction collector (Agilent™, Santa Clara, CA, USA). The samples were loaded in the column equilibrated with 60% solvent A (60:40 acetonitrile:water + 10mM ammonium acetate) in 40% solvent B (90:10 isopropanol:acetonitrile + 10mM ammonium acetate) and the elution was performed at a flow rate of 0.5 mL/min using the following gradient: 40-50%B in 2 min, 50-60% B in 1 min, 60-70% B in 9 min, hold in 70% B for 3 min, 70-78% B in 2 min, 78-85% B in 2 min, 85-99%B in 6 min and hold at 99% B for 3 min. Eluting lipids were monitored by electrospray ionization mass spectrometry (LTQ™ Linear Ion Trap Mass Spectrometer, Thermo Scientific™, Waltham, MA, USA) with 26 fractions collected in regular intervals of 1 min from RT 2.5 min – 28.5 min. Fractions from 18 fractionation runs were dried in a vacuum centrifuge and dissolved in appropriate solvent for subsequent analyses.

**Proteomic analysis**

Extracted proteins were suspended in 100 µL of 8 M urea in 50 mM NH₄HCO₃ and protein concentration was determined by a BCA protein assay (Thermo Scientific, Rockford, IL). Samples were digested as previously described (54) and desalted using 1 mL Discovery C18 SPE columns (Supelco, Bellefonte, PA). Digested peptides were suspended in water, quantified by BCA assay and had their concentrations adjusted to 0.1 µg/µL with water. A total of 500 ng of peptides were direct injected onto the analytical column. The analytical column used was prepared in-house by slurry packing 3-µm Jupiter C18 stationary phase (Phenomenex, Torrence, CA) into a 70-cm long, 360 µm OD x 75 µm ID fused silica capillary tubing (Polymicro Technologies Inc., Phoenix, AZ). The column was coupled to a Waters M-Class™ UPLC system (Millford, MA) connected with a Q-Exactive Plus hybrid quadrupole/Orbitrap mass spectrometer from Thermo Fisher Scientific (San Jose, CA). Mobile phases consisted of 0.1% formic acid in water (MP-A) and 0.1% formic acid in acetonitrile (MP-B). The gradient elution, and column regeneration profile was as follows (min:%MP-B); 0:0.1, 4.0:8, 36:12, 135:30, 175:45, 180:95, 190:95, 195:1. Data acquisition (180 min) was started at the end of the sample loading period (20 min). The analytical column was coupled to the Q-Exactive using a home-built nanospray adapter interface with 2.2 kV applied to achieve electrospray ionization. The MS inlet was maintained at a temperature of 275 °C. A precursor scan was performed from m/z 400 to 2000 at a resolution of 35k at m/z 400 and an automatic gain control (AGC) of 3e6. Operated in data dependent mode, the top 12 most intense ions from the precursor scan were selected for high energy collision dissociation (HCD) MS/MS at a resolution of 17.5k, AGC of 1e5, isolation window of 2 m/z, and a max ion time of 100 ms. Only ions identified as having a +2
charge or higher were subjected to HCD and subsequently excluded from further analysis for 30 sec thereby allowing for deeper coverage.

LC-MS/MS data were analyzed with MaxQuant software (v.1.5.5.1) by searching against the *H. capsulatum* G186AR (downloaded August 15, 2016) and *S. cerevisiae* S288c (downloaded January 11, 2018) sequences from Uniprot Knowledgebase, considering protein N-terminal acetylation and oxidation of methionine as variable modifications, and cysteine carbamidomethylation as a fixed modification. The parent ion mass tolerance was set at 20 and 4.5 ppm for the first and main peptide searches, whereas the fragment mass tolerance was set as 20 ppm for both searching rounds. Tryptic digestion was required for peptide termini and up to 2 missed cleavage sites were allowed per peptide. Protein abundances were estimated by the intensity based absolute quantification (iBAQ) method (57). The intensities were normalized by the total iBAQ sum of each sample and expressed as relative number of protein copies (percentage from total). Proteins were classified according to their abundance converting number of copies described by Beck et al. (29) to relative number of protein copies (%). Function-enrichment analysis was performed as described previously (58).

**Lipid analysis by liquid chromatography-tandem mass spectrometry**

Total lipid extracts and phospholipid fraction from the Silica 60 column were resuspended in methanol and subjected to LC-MS/MS analysis as previously described (59). To assess the technical variability, we spiked in the SPLASH Lipidomix standard (Avanti Polar), which contains a mix isotopically labeled lipids. Lipid species were identified using LIQUID and manually inspected for validation. The features of the identified lipid species were extracted and aligned using MZmine (60). For comparative purposes, lipids were considered significantly different with \( p \leq 0.05 \) by \( t \)-test considering equal variance and unpaired samples. The distribution of fatty acyl groups was done by counting individual fatty acyl groups and were considered significantly different with a \( p \leq 0.05 \) by Fisher’s exact test.

**Lipid Analysis by ion mobility spectrometry-mass spectrometry (IMS-MS)**

HPLC-fractionated fractions were suspended in methanol and analyzed on an Agilent 6560 IMS-QTOF MS platform (Agilent Technologies, Santa Clara, CA) (61). For IMS measurements, ions were generated by an electrospray ionization source, passed through the glass capillary, focused and accumulated in the ion funnel before being pulsed into the IMS drift tube filled with \( \sim 3.95 \) torr of nitrogen gas, where they traveled under the influence of a weak electric field (10-20 V/cm). Ions exiting the drift tube are then refocused by a rear ion funnel prior
to being detected by QTOF MS, and their arrival times were recorded. Collision-cross section (CCS) which contains information of molecule size and shape can be determined from the data acquired under different electric fields. The Agilent IM-MS Browser software was used to process IMS data, extract IMS profiles and calculate CCS values. Candidate phospholipid species were matched to standards based on their CCS values and MS/MS spectra.

**Fatty acid and sterol analyses**

Fatty acid fractions from the silica 60 column fractionation were resuspended in 0.5 mL of anhydrous methanolic HCl (1.2 N), sealed and incubated for 1 h at 100°C. After cooling, 0.5 mL of water was added and the fatty acid methyl esters (FAMEs) were extracted by adding 0.5 mL hexane and vortexing the samples. Sterol fractions were derivatized with N-methyl-N-(trimethylsilyl)trifluoroacetamide (MSTFA) (Sigma-Aldrich) and trimethylchlorosilane (TMCS) (Sigma-Aldrich) as described in details elsewhere(62). Derivatized FAMEs and sterols were then analyzed by GC-MS. For the background referencing and retention time calibration, blanks and FAME standard mix (C8-28) (Sigma-Aldrich) were also analyzed. Analyses were performed in an Agilent GC 7890A using a HP-5MS column (30 m × 0.25 mm × 0.25 μm; Agilent Technologies, Santa Clara, CA) coupled with a single quadrupole MSD 5975C (Agilent Technologies). For each analysis 1 μL of sample was injected (splitless) into the injection port with constant temperature at 250°C. The GC analysis started with an initial oven temperature fixed at 60°C for 1 minute after injection, and then the temperature was increased to 325°C at a rate of 10°C/minute, and finished with a 5-minute hold at 325°C. The data files were calibrated and deconvoluted with Metabolite Detector as stated previously(63). Molecules were identified by library matching against the FiehnLib (64) with some in-house added entries. Unidentified chromatographic features were also searched against the Wiley Fatty Acids Methyl Esters Mass Spectral Database and the NIST17 GC-MS spectral library.

**Construction of the *H. capsulatum* lipid metabolic map**

We have built a preliminary G217B lipid map with a tool name VANTED v2.6.5(65), using the *S. cerevisiae* model (30) as a starting point. Enzyme orthologues were identified by blasting *S. cerevisiae* and *C. neoformans*, two of the best characterized fungi in terms of lipid metabolism (30, 66). The map was integrated with abundance level of lipids using their relative mass spectrometry intensities within the same lipid class.
Evaluation of lipid biosynthesis pathways on *H. capsulatum* growth

The tested compounds (myriocin, 10-thiastearic acid and thiocarlide) were serially diluted in round bottom 96-well plates. *H. capsulatum* yeast cells were washed with PBS and suspended in Ham’s F12. Yeast cells were added in 4 replicates to the drug-containing plates at a final cellular density of 2.5 x 10⁴ cells per mL. The plates were incubated at 37 °C under constant shaking for 7 days until the optical analysis of growth. The drug concentration present in the set of replicates with no visual growth was considered the MIC for each of the drugs. To evaluate the effect of lipid synthesis inhibition of *H. capsulatum*, 10⁶ cells/mL were incubated in Ham’s F12 in the presence or absence of myriocin, 10-thiasteric acid and thiocarlide in half of the evaluated MIC for each drug. The cultures were kept at 37°C shaking at 250 rpm for 48 hours. Cells were extensively washed with ice-cold PBS and pellets were frozen for further lipid extraction and subsequent fatty acid and lipidomics analyses. This experiment was performed in triplicates.

Platelet aggregation assay

Platelet-rich and platelet-depleted plasma were obtained from human blood by differential centrifugation. Platelet aggregation was measured in a Chrono-Log 700 aggregometer by impedance in whole blood.

Treatment of macrophages with *H. capsulatum* lipid fractions

J774 macrophage-like cell line was cultivated in DMEM medium containing 10% FBS at 37°C and 5% CO₂ atmosphere. Cells were then treated for 24 hours with lipids fractionated by HPLC and resuspended in methanol. Due to the lack of the exact concentration of the lipid fractions, these were tested in 4 dilutions: 10⁻³, 10⁻⁴, 10⁻⁵ and 10⁻⁶. Control treatments were carried in the presence or absence of the PAF receptor antagonist WEB 2086, and vehicle only (same amount as in the highest concentration of the fractions – 0.1%). After treatment TNF-α and IL-10 were measured by ELISA (B&D). Results were analyzed with GraphPad (Prism) and were considered significant with p ≤ 0.05 by ANOVA followed by Tukey’s multiple comparison test.
AUTHOR CONTRIBUTIONS
D.Z.M., H.M.H., M.C.B., J.D.N and E.S.N. designed the research. D.Z.M. performed the H. capsulatum growth, drug testing and cytokine experiments. H.M.H., M.C.B., E.M.Z., J.E.K., K.K.W., E.L.B. and K.J.B. performed the lipidomics and proteomics experiments. E.L.B. performed the genetic engineering of the S. cerevisiae strains. M.C.B., H.M.H., N.M.M. and Y.M.K. performed the gas chromatography-mass spectrometry analyses. E.S.N., G.C., J.D.Z., J.R.T. build the lipid metabolic map. X.Z. and E.S.B. performed the ion mobility-mass spectrometry analysis. M.R.G. performed the platelet aggregation analysis. D.Z.M., H.M.H., M.C.B., S.P.C., X.Z., E.S.B, J.E.K., S.H.P., Y.M.K., M.R.G., J.D.N., and E.S.N. analyzed the data. M.R.G., E.S.B, S.H.P., J.D.N and E.S.N. contributed with reagents and resources. D.Z.M., H.M.H, M.C.B. and E.S.N. wrote the manuscript with inputs from the other authors. All the authors read and approved the final version of the manuscript.

CONFLICTS OF INTEREST
The authors declare no financial conflicts of interest.

Acknowledgments
The authors thank Drs. Igor Almeida, Rosa Maldonado, Milene Carmes Vallejo, Charles Ansong and Joshua Adkins for insightful discussions. Joshua D. Nosanchuk and Ernesto S. Nakayasu were supported by NIH R21 AI124797. Erin L. Bredeweg, Ernesto S. Nakayasu, Jennifer Kyle were supported by a Laboratory Directed Research and Development project from Pacific Northwest National Laboratory (PNNL). Erin S. Baker would also like to acknowledge support from NIEHS P42 ES027704. Parts of this work were performed in the Environmental Molecular Science Laboratory, a U.S. Department of Energy (DOE) national scientific user facility at PNNL in Richland, WA.

Data availability. Proteomics data were deposited into Pride repository (www.ebi.ac.uk/pride) under accession number PXD017734.
Username: reviewer86800@ebi.ac.uk
Password: 9cZs0O1G
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Figure legends

**Fig. 1** | Proteomic and lipidomic analyses of *Histoplasma capsulatum* yeast cells. (a) Extraction procedure for lipidomics proteomics analyses. Yeast cells were extracted sequentially with chloroform:methanol (2:1, v:v) and chloroform:methanol:water (1:2:0.8, v:v:) and fractionated by Silica 60 solid phase extraction for sterol, free fatty acid and phospholipid analyses. Cells were also submitted to simultaneous metabolite, protein and lipid extraction (MPLEx) for global lipidomics and proteomics analyses. (b) Overall lipid coverage combining MPLEx and specific lipid extractions of *Histoplasma capsulatum*. (c) Protein abundance classification based on their relative copy numbers. (d) Function enrichment analysis based on the KEGG annotation of *H. capsulatum* proteins with different abundance levels. Abbreviations: C, chloroform; Cer, ceramide; CL, cardiolipin; DG, diacylglycerol; FA, fatty acid; HexCer, hexosylceramide; LPC, lysophosphatidylcholine; LPE, lysophosphatidylethanolamines, LPG, lysophosphatidylglycerol; LPI, lysophosphatidylinositol; LPS, lysophosphatidylserine; M, methanol; PA, phosphatic acid; PC, phosphatidylinositol; PE, phosphatidylethanolamine; PG, phosphatidylglycerol; PI, phosphatidylinositol; PI_Cer, inositolphosphoceramide; PL, phospholipid; PS, phosphatidylserine; ST, sterol; TG, triacylglycerol; W, water.

**Fig. 2** | Integrated metabolic map of *Histoplasma capsulatum* lipids and biosynthesis proteins. The map was built based on the genomic information along with proteomics and lipidomics data. Lipids abundances were normalized by the most intense signal in the mass spectrometry analysis within each lipid class, whereas the proteins were classified based on their relative copy number. For details about enzyme names and orthologues see table S10. Abbreviations: Cer, ceramide; CL, cardiolipin; DG, diacylglycerol; HexCer, hexosylceramide; LPC, lysophosphatidylcholine; LPE, lysophosphatidylethanolamines, LPG, lysophosphatidylglycerol; LPI, lysophosphatidylinositol; LPS, lysophosphatidylserine; M, methanol; PA, phosphatic acid; PC, phosphatidylinositol; PE, phosphatidylethanolamine; PG, phosphatidylglycerol; PI, phosphatidylinositol; PI_Cer, inositolphosphoceramide; PL, phospholipid; PS, phosphatidylserine; ST, sterol; TG, triacylglycerol.

**Fig. 3** | Analysis of *Saccharomyces cerevisiae* (Sc) and *Histoplasma capsulatum* (Hc) lysophospholipid acetyltransferase LPT1 genes. The analysis was done in wild-type and LPT1-knockout (KO) (*Lpt1* -/-) *S. cerevisiae* strains complemented or not with the pESC-URA plasmid with Hc-LPT1 gene under galactose-inducible promoter. *S. cerevisiae* WT and LPT1-knockout (KO) strains were grown in YNB medium supplemented with of glucose (Glc) or galactose (Gal). (a) Protein abundance of Sc and Hc LPT1 proteins measured by the intensity of specific peptides in the LC-MS/MS analysis. Elongation factor Tu (*TUF1*) was used as a loading control. As expected, Sc LPT1 was only detected in the WT strain, whereas Hc LPT1 was detected only in strains transformed with the plasmid and induced with galactose. (b-c) Heatmaps of identified HcLPT1 (b) and ScLPT1 (c) products (p≤0.05 compared the complemented vs control strains (b) or LPT1-KO vs. WT strains (c). The experiments were done in two independent batches delimited by the vertical grey lines. (d-e) Distribution of the lipid classes (d) and fatty acyl groups (e) of the HcLPT1 and ScLPT1 products. *p≤0.05 by Fisher's exact test.
**Fig. 4** | **Fatty acid metabolism in *Histoplasma capsulatum***. (a) Distribution of carbon chain lengths of fatty acyl groups from lipids in global lipidomic analysis. (b) Fatty acid desaturase pathways in *H. capsulatum*, *Saccharomyces cerevisiae* and Human. (c) Distribution of double bonds on fatty acyl groups from lipids in global lipidomic analysis. (d) Minimum inhibitory concentrations (MIC) of fatty acid desaturase inhibitors. The MICs were obtained from 5 independent experiments with 4 replicates each. (e) Effect of fatty acid desaturase inhibitors in the fatty acid profile. Cells were grown in one half of each compound MIC and submitted to fatty acid analysis. The values represent the ratio of peak areas of unsaturated/saturate fatty acids. Statistical tests: *p≤0.05 by Fisher’s exact test and # p≤0.05 by Student t-test.

**Fig. 5** | **Sphingolipid metabolism in *Histoplasma capsulatum***. (a) Sphingolipids of *H. capsulatum*, *Saccharomyces cerevisiae* and Human. (b) Minimum inhibitory concentration (MIC) of myriocin, the inhibitor of serine-palmitoyl transferase, which is the first step of the sphingolipid biosynthesis. The MICs were obtained from 5 independent experiments with 4 replicates each. (c) Effect of myriocin on the lipidomics profile. Cells were grown in one half of myriocin MIC and submitted to fatty acid analysis. The values represent the mass spectral signal intensity on the apex of the peak. Statistical tests: *p≤0.05 by Fisher’s exact test and # p≤0.05 by student t-test.

**Fig. 6** | **Identification and validation of platelet-activating factor (PAF) analogs in *Histoplasma capsulatum***. (a) Structure, monoisotopic mass and tandem mass (MS/MS) fragments of PAF and analogs. (b) LC-MS/MS analysis of *H. capsulatum* PC (16:0/2:0) against the synthetic standard. The graphs showed the same retention time of extracted ion chromatograms of *H. capsulatum* phospholipids, *H. capsulatum* phospholipids + standard and standard alone. (c-d) MS/MS spectrum (c) and ion mobility-mass spectrometry (IMS-MS) (d) analyses of the PC (16:0/2:0) compared to the synthetic standard. The MS/MS analysis show the diagnostic fragment of phosphatidylcholines at m/z 184.07 (d), whereas the IMS-MS shows that the molecule has compatible arrival time and collision cross sections (stated in Å²) with the standard.

**Fig. 7** | **Effect of fractions enriched in *Histoplasma capsulatum* PAF analogs in platelet aggregation**. Human peripheral blood was incubated with the indicated positive controls and fractions enriched in *H. capsulatum* PAF analogs and aggregation was measured under an aggregometer. Panels a and b are two measurements of platelet aggregation activity of HPLC fractions enriched with *H. capsulatum* PAF analogs (fractions 8-10). The arrows indicate the addition of each compound or HPLC fraction. The figure is representative of two independent experiments with similar results.

**Fig. 8** | **Cytokine production by J774 macrophages stimulated with HPLC fractions enriched in *Histoplasma capsulatum* PAF analogs**. J774 macrophages were pre-treated or not with the PAF receptor antagonist (WEB2086 – 10 mM) for 30 minutes before the incubation with acyl-PAF PC (16:0/2:0) and HPLC fractions enriched in *H. capsulatum* PAF analogs (fractions 9-10) for 24 hours. After the incubation period, TNF-α (a) and IL-10 (b) were quantified from the cell culture supernatants by ELISA. The graphs show averages and SEM for 3 experiments. The analysis reviewed an induction of cytokine production by HPLC fractions.
enriched in *H. capsulatum* PAF analogs, which were inhibited by the PAF receptor agonist WEB2086 (WEB). *p < 0.05* for groups compared to the solvent control (first bar). +p < 0.05 for respective groups in the presence and absence of WEB2086 by ANOVA followed by Tukey's multiple comparison test.
Supplementary fig. 1 | Tandem-mass (MS/MS) spectra of the platelet-activating factor analog PC(16:2/2:0). Abbreviations: Ac, acetate; Cho, choline; ChoP, phosphocholine; ChoPGlyc, glycerophosphocholine; CID, collision-induced dissociation; DG, diacylglycerol; HCD, high-energy induced dissociation; M, mass; m/z, mass-over-charge ratio; ppm, parts-per-million. The spectra show diagnostic fragments for phosphatidylcholine (Cho, ChoP and ChoPGlyc) and for an acetyl group (M-Ac-H2O).

Supplementary fig. 2 | Tandem-mass (MS/MS) spectra of the platelet-activating factor analog PC(16:1/2:0). Abbreviations: Ac, acetate; Cho, choline; ChoP, phosphocholine; ChoPGlyc, glycerophosphocholine; CID, collision-induced dissociation; DG, diacylglycerol; HCD, high-energy induced dissociation; LPA, lysophosphatidic acid; M, mass; m/z, mass-over-charge ratio; ppm, parts-per-million. The spectra show diagnostic fragments for phosphatidylcholine (Cho, ChoP and ChoPGlyc) and for an acetyl group (M-Ac-H2O and LPA(16:1)-H2O).

Supplementary fig. 3 | Quantification of PC(16:1/2:0) in wild-type (WT) and Lpt1-knockout (KO) Saccharomyces cerevisiae strains transformed with a plasmid containing the Histoplasma capsulatum Lpt1 gene (pHcLPT1). S. cerevisiae strains complemented or not with the pESC-URA plasmid with Hc-LPT1 gene under galactose-inducible promoter and grown in YNB medium supplemented with galactose. The experiment was performed with 4 replicates. Statistical test was done with student t-test considering equal distribution.

Supplementary tab. 1 | List of identified lipids in the global lipidomics analysis.

Supplementary tab. 2 | Quantification of identified lipids in the positive mode of global lipidomics analysis. The quantitative information is based on the extracted intensity of each lipid species at the peak apex. The measurements were performed in 4 replicates.

Supplementary tab. 3 | Quantification of identified lipids in the positive mode of global lipidomics analysis. The quantitative information is based on the extracted intensity of each lipid species at the peak apex. The measurements were performed in 4 replicates.

Supplementary tab. 4 | List of identified lipids in the phospholipidomics analysis.

Supplementary tab. 5 | Quantification of identified lipids in the positive mode of phospholipidomics analysis. The quantitative information is based on the extracted intensity of each lipid species at the peak apex. The measurements were performed in 4 replicates.

Supplementary tab. 6 | Quantification of identified lipids in the positive mode of phospholipidomics analysis. The quantitative information is based on the extracted intensity of each lipid species at the peak apex. The measurements were performed in 4 replicates.
Supplementary tab. 7 | Quantification of identified sterol species in the gas chromatography-mass spectrometry analysis. The quantitative information is based on the extracted peak areas of each lipid species. The measurements were performed in 4 replicates.

Supplementary tab. 8 | Quantification of identified fatty acid species in the gas chromatography-mass spectrometry analysis. The quantitative information is based on the extracted peak areas of each lipid species. The measurements were performed in 4 replicates.

Supplementary tab. 9 | Quantification of identified proteins in the proteomics analysis. The quantitative information is based on the intensity based absolute quantification method (iBAQ), which was further converted to relative copy number of proteins per cell.

Supplementary tab. 10 | List of orthologues from *Saccharomyces cerevisiae* S288c and *Cryptococcus neoformans* H99 to *Histoplasma capsulatum* G186AR.

Supplementary tab. 11 | Lipid species upregulated in *Saccharomyces cerevisiae* strains expressing the *Histoplasma capsulatum* Lpt1 gene. The analysis was done in wild-type (WT) and LPT1-knockout (KO) *S. cerevisiae* strains complemented or not with the pESC-URA plasmid with Hc-LPT1 gene under galactose-inducible promoter. *S. cerevisiae* WT and LPT1-knockout strains were grown in YNB medium supplemented with glucose (Glc) or galactose (Gal). The measurements were performed in 4 replicates. Statistical analysis was done with student *t*-test considering equal distribution.

Supplementary tab. 12 | Lipid species downregulated in *Saccharomyces cerevisiae* strains expressing the *Histoplasma capsulatum* Lpt1 gene. The analysis was done in wild-type (WT) and LPT1-knockout (KO) *S. cerevisiae* strains complemented or not with the pESC-URA plasmid with Hc-LPT1 gene under galactose-inducible promoter. *S. cerevisiae* WT and LPT1-knockout strains were grown in YNB medium supplemented with glucose (Glc) or galactose (Gal). The measurements were performed in 4 replicates. Statistical analysis was done with student *t*-test considering equal distribution.

Supplementary tab. 13 | Lipid species downregulated in *Saccharomyces cerevisiae* Lpt1 knockout strain. The analysis was done in wild-type and LPT1-knockout (KO) strains grown in YNB medium supplemented with glucose (Glc) or galactose (Gal). The measurements were performed in 4 replicates. Statistical analysis was done with student *t*-test considering equal distribution.
**Fig. 1**

**a**

H. capsulatum cells

C:M (2:1)  
C:M:W (1:2:0.8)  
MPEX  
Silica 60  
Lipidomics  
Trypsin digestion

ST  
FA  
PL  
Proteomics

**b**

Glycerolipid (149)

Sterols (6)  
Fatty acids (9)  
Sphingolipids (12)

Phospholipids (171)

Lyso-phospholipids (24)

**c**

Relative copy number (%)

Abundance rank

**d**

| Pathway                                      | High | Moderate | Low | Very Low |
|----------------------------------------------|------|----------|-----|----------|
| Glycolysis / Gluconeogenesis                 | 5.9  | 1.1      | 0.0 | 0.0      |
| Citrate cycle (TCA cycle)                    | 8.7  | 0.0      | 0.0 | 0.0      |
| DNA replication                              | 1.3  | 3.9      | 2.4 | 2.3      |
| Nicotinate and nicotinamide metabolism       | 0.6  | 1.7      | 4.1 | 0.0      |
| Basal transcription factors                  | 0.3  | 2.4      | 4.4 | 7.0      |
| Fatty acid metabolism                        | 4.5  | 1.6      | 0.4 | 0.0      |
| Fatty acid biosynthesis                       | 3.8  | 1.4      | 0.0 | 0.0      |
| Fatty acid elongation                        | 4.7  | 1.8      | 1.5 | 0.0      |
| Fatty acid degradation                       | 5.5  | 1.2      | 0.0 | 0.0      |
| Arachidonic acid metabolism                  | 4.7  | 0.0      | 0.0 | 0.0      |
| alpha-Linolenic acid metabolism              | 1.6  | 1.2      | 0.0 | 0.0      |
| Biosynthesis of unsaturated fatty acids      | 3.1  | 1.8      | 0.0 | 0.0      |
| Steroid biosynthesis                         | 1.7  | 2.9      | 1.6 | 0.0      |
| Glycerophospholipid metabolism               | 0.5  | 2.2      | 2.7 | 3.4      |
| Ether lipid metabolism                       | 0.9  | 0.6      | 6.4 | 0.0      |
| Sphingolipid metabolism                      | 1.4  | 2.2      | 1.8 | 0.0      |
| Glycosylphosphatidylinositol (GPI)-anchor biosynthesis | 0.0  | 1.5      | 1.2 | 0.0      |
Fig. 3
Fig. 4
a

\textbf{H. capsulatum}  
Ceramide  
\[ \begin{array}{c}
\text{Glc-Cer} \\
\downarrow \\
\text{PI-Cer} \\
\downarrow \\
\text{Man-PI-Cer} \\
\downarrow \\
\text{Gal-Man}_2\text{-PI-Cer}
\end{array} \]

\textbf{S. cerevisiae}  
Ceramide  
\[ \begin{array}{c}
\text{Cer-OH} \\
\downarrow \\
\text{PI-Cer} \\
\downarrow \\
\text{Man-PI-Cer} \\
\downarrow \\
\text{Man-(PI)}_2\text{-Cer}
\end{array} \]

\textbf{Human}  
Ceramide  
\[ \begin{array}{c}
\text{SM} \\
\downarrow \\
\text{Globo} \\
\downarrow \\
\text{Lacto} \\
\downarrow \\
\text{Isoglobo}
\end{array} \]

b

\begin{tabular}{lc}
\textbf{Drug} & \textbf{MIC (μM)} \\
\hline
Myriocin & 0.03 \\
\hline
\end{tabular}

Fig. 5
Fig. 6
Fig. 7
Fig. 8
PC(16:2/2:0)
Mass error = -1.75 ppm

High resolution - HCD

Low resolution - CID

Supplementary fig. 1
PC(16:1/2:0)
Mass error = -2.00 ppm

High resolution - HCD

[ChoP]^+ 184.07
[Cho]^+ 104.11

[ChoPGlyc]^+ 258.11
[DG-H2O+H]^+ 353.27

[M-H2O]^+ 518.32

[M-Ac-H2O]^+ 476.28

Relative Abundance

Supplementary fig. 2.
Supplementary fig. 3.