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Article

A Comprehensive Analysis of the Lipidomic Signatures in *Rhizopus delemar*

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Abstract: Certain species of Mucorales have been identified as causative agents of mucormycosis, a rare yet often lethal fungal infection. Notably, these fungi exhibit intrinsic resistance to common azole drugs, which target lipids. Given the pivotal role of lipids in drug resistance and their contribution to the innate resistance to azoles, this study provides a comprehensive overview of key lipid classes, including sphingolipids (SLs), glycerophospholipids (GPLs), and sterols, in *Rhizopus delemar* 99-880, a well-characterized reference strain among Mucorales. Through high-throughput analysis involving the identification and quantification of intermediates, we have elucidated the biosynthetic pathways of SLs, PGLs, and sterols. Notably, the acidic SLs were not found probably because the acidic branch of the SL biosynthesis pathway terminates at α -hydroxy phytoceramides as evident by their high abundance. Intermediates in the neutral SL pathway incorporated higher levels of 16:0 fatty acid compared to other pathogenic fungi. A strikingly high phosphatidylethanolamine (PE)/phosphatidylcholine (PC) ratio was observed among GPLs. Ergosterol remains the major sterol, similar to other fungi, and our analysis confirms the existence of alternate ergosterol biosynthesis pathways. The total lipidomic profile of *R. delemar* 99-880 offers insights into its lipid metabolism and potential implications for studying pathogenesis and drug resistance mechanisms.

Keywords: mucormycosis; drug resistance; posaconazole; amphotericin B; sphingolipids; phospholipids; sterols; mass-spectrometry

1. Introduction.

Mucormycosis is a serious disease caused by a group of fungi called Mucorales. There are more than two dozen species of these fungi that can cause this deadly disease. The most common culprits are species from the genera *Rhizopus* and *Mucor*, with *Rhizopus arrhizus* being the most prevalent [1]. People with diabetes, neutropenia, and those who have received hematopoietic or solid organ transplants are more susceptible to mucormycosis. Unfortunately, the number of diabetic patients, especially in developing countries, is increasing, which puts them at a higher risk for this disease. In fact, diabetes is a major risk factor for mucormycosis in over 70% of cases in countries like India, compared to 36-40% globally [2-4].

Mucormycosis is challenging because treatment with first-line therapy of lipid formulation amphotericin B formulations are often associated with severe toxicities that in many cases limit their use [5,6]. Importantly, Mucorales fungi exhibit higher MICs and in many cases, resistance to azoles (e.g. posaconazole [POSA] and isavuconazole [ISAV]) that are used as a stepdown therapy for

mucormycosis [7,8]. The disease is intrinsically resistant to short tailed azoles like fluconazole (FLC) and voriconazole (VCZ) which otherwise are the most widely used antifungal drugs[9].

Studies on azole resistance mechanisms in various pathogenic fungi have revealed similarities across different species. These mechanisms often involve the overexpression of efflux pumps, mutations in target gene *ERG11*, or increased expression of target enzyme lanosterol 14 α -demethylase (LDM) [10,11]. It is important to note that the prevalence of these resistance mechanisms can vary among different fungi. For example, in *Candida* species, the most common mechanism is the overexpression of efflux pump genes, while in *Aspergillus fumigatus*, mutations in the azole target enzyme lanosterol 14 α -demethylase (LDM) are more prevalent [12]. *Cryptococcus* species have been found to exhibit overexpression of various genes due to reversible chromosomal duplications, point mutations in *ERG11*, and increased efflux pump activity [13,14]. In contrast, the molecular basis of drug resistance in Mucorales is not well understood and remains an area that requires further exploration. Studies have shown that the genome of *Mucor circinelloides* contains eight potential PDR-type transporters, and the regulation of these genes appears to be interconnected. Both Pdr1 and Pdr2 have been found to contribute to the resistance of *M. circinelloides* against certain azole drugs like PCZ, ICZ, and ravuconazole (RVZ). However, it is worth noting that azole resistance in general cannot be fully explained solely by the activity of these tested efflux pumps. The involvement of other PDR genes and novel mechanisms cannot be ruled out and may play a role in drug resistance [7].

Azoles represent a crucial class of medications targeting lipids, specifically lanosterol 14 α -demethylase (LDM), an enzyme encoded by the *ERG11* gene. By inhibiting LDM, azole drugs impede the production of ergosterol, a vital component of fungal cell membranes, resulting in the build-up of toxic sterols [15]. While our comprehension of acquired resistance to triazole drugs in fungi like *Aspergillus* and *Candida* species is substantial, the inherent resistance of Mucorales to short-tailed triazoles such as VCZ and FLC warrants further exploration [16]. In many fungal species, acquired azole resistance often arises from specific mutations in the LDM gene, reducing affinity for certain triazole drugs. However, the absence of *in vitro* activity of FLC and VCZ against Mucorales suggests their resistance may be innate rather than acquired, possibly due to ancient amino acid substitutions. A 2017 study by Rita Caramalho *et al.* aligned LDM sequences from six Mucorales species, identifying a conserved Y129F substitution likely responsible for innate resistance to short-tailed azoles like FLC and VCZ [17]. Analysis, based on the known LDM structure, indicates a mechanism for this mutation involving the disruption of a water-mediated hydrogen bond [17]. Significantly, introducing this mutation into the homologous protein of *Saccharomyces cerevisiae* conferred resistance to short-tailed azoles [18]. Other point mutations in genes involved in ergosterol biosynthesis have also been observed in Mucorales, raising concerns about the implications of emerging resistance to current antifungal drugs. Further research is needed to fully understand the mechanisms of innate resistance in Mucorales and to develop effective strategies for combating this resistance.

Research has highlighted the significant role of lipids, the target of common antifungal drugs, in determining drug susceptibility and virulence in various pathogenic fungi [19]. However, this aspect remains poorly understood in Mucorales. Considering the severity of Mucorales infections as secondary complications in Covid-19 and diabetes patients [20], it becomes imperative to gain a comprehensive understanding of the lipid landscape and imbalances to facilitate the development of effective treatments. In this study, we selected the genome-sequenced *Rhizopus delemar* 99-880 strain as a reference strain to investigate the lipidomic landscape of Mucorales [21]. Utilizing high-throughput liquid chromatography electro spray ionisation mass spectrometry (LC-ESI/MSMS) and gas chromatography coupled to mass spectrometry (GCMS) approaches, we successfully conducted qualitative and quantitative analyses of a wide range of lipids including sphingolipids (SLs), glycerophospholipids (GPLs), and sterols. This dataset presents, for the first time, a comprehensive overview of the lipidomic profile in the well-defined *R. delemar* 99-880 strain, which is anticipated to serve as a foundational platform for future investigations in this field.

2. Materials and Methods.

2.1. Strains, Media and Growth Conditions.

R. delemar 99-880 spores were revived from archived glycerol stocks stored at -80°C. Spores were inoculated on YPD plates and incubated for 3-4 days until spores were formed. Plates were flooded with 10 mL phosphate buffered saline with 0.01% tween (PBST) and spores were collected by aspiration. Spores were counted using hemocytometer and adjusted to a density of 10^6 spores mL⁻¹ for further experimental use.

2.2. Drug Susceptibility.

Approximately 10^3 spores were spotted on YPD plates containing different concentrations of the drugs, Myriocin (MYR) (Sigma) or Aureobasidin A (AbA) (Takara). Plates were incubated at 30°C for 48 hours and then imaged (Bio-rad Molecular Imager®).

2.3. Sphingolipid Isolation.

Spores were inoculated in 100 mL YPD and grown at 30°C with constant shaking until mycelia were formed (~24 hours). Mycelia were harvested and washed with 0.9% saline and dried. Approximately 100 mg of mycelia were used for SL isolation using methods as described earlier by Kumar *et al* [22]. C17 sphingosine and C17 ceramide (Avanti Polar Lipids Inc. AL, USA) were added as internal standards before isolation. Dried SL extracts were dissolved in 300 μ L organic buffer (methanol with 1 mM ammonium formate and 0.2% formic acid). From this, 20 μ L sample was transferred to glass insert and final volume adjusted to 200 μ L in organic buffer for LC-MS/MS analysis.

2.4. Phospholipid Isolation.

PLs were extracted using the Folch method with slight modifications[23]. Equisplash® (Avanti Polar Lipids Inc. AL, USA) containing an equal proportion of deuterated PLs of each class was added as internal standard before isolation. Briefly 100 mg mycelia were homogenized in 1 mL LC/MS grade water using glass beads in FastPrep(MP Biomedicals). To this lysate, 9 mL of 2:1 chloroform:methanol was added and incubated for 2 hours with periodic vortexing. The upper layer was discarded and 2 mL of 0.9% saline was added. Lower layer was then separated by centrifugation at 3000 rpm for 5 min and dried with N₂ gas. Dried extracts were dissolved in 1 mL organic buffer. From this 10 μ L sample was transferred to glass insert and final volume adjusted to 200 μ L in organic buffer for LC-MS/MS analysis.

2.5. Sterol Extraction.

Harvested mycelia were finely crushed in liquid N₂ and approx. 20 mg of it was weighed and used for sterol extraction. Sterols were extracted as described by Morio *et al* [24]. Mycelia was resuspended in 300 μ L saponification solution containing 12.5g KOH dissolved in 18 mL MQ water and adjusted to 50 mL with 98% ethanol. The mixture was then heated at 80°C for 1h in a capped glass vial. After cooling at RT, 100 μ L MQ water, 400 μ L hexane and 1 μ L cholestane (5 mg/ml) as internal standard was added. The mixture was vortexed and after phase separation, the top layer of ~350 μ L (hexane) was separated and 600 μ L hexane was again added to the remaining lysate. After vortexing and phase separation, 550 μ L of the top layer was separated and both hexane fractions were combined and dried in vacuum centrifugation at room temperature. The dried sterol extracts were re-dissolved in 60 μ L hexane and 10 μ L Silylation mixture (Sigma, 85432) for derivatization, vortexed and incubated for 1 hour at room temperature. From this 50 μ L of the debris-free extract was transferred into glass inserts for GCMS analysis.

2.6. Liquid Chromatography-Mass Spectrometry.

SLs were analyzed by employing Shot-gun and multiple reaction monitoring (MRM) approaches using LC-MS (QTRAP® 4500, AB SCIEX). For untargeted shotgun lipidomics, different confirmatory scans were used and the spectra obtained are given in **Supplementary File S1**. For the targetted MRM approach, SLs and PLs were separated on C8-Column and C18 columns maintained at 60°C and 50°C

respectively (Waters, USA). Organic buffer (methanol with 1 mM ammonium formate and 0.2% formic acid) and aqueous buffer (water with 2 mM ammonium formate and 0.2% formic acid) were used as mobile phase. A combined flow rate of 300 μ L per min was maintained in a gradient manner starting with 80% organic buffer, gradually increased to 88%, then 99%, decreased to 88% and finally restored to 80% before final run. Mass spectrometric parameters were standardized as follows: Source temperature was set at 600°C for SL, nebulizer and desolvation gases (GS1 and GS2 respectively) were set at 50 psi each, electrospray ionization voltage at 5500 V and curtain gas at 45 psi. For PL analysis the source temperature was set as 400°C, GS1 and GS2 at 45 psi, electrospray ionization voltage at 4500 V and curtain gas at 45 psi.

2.7. Gas Chromatography-Mass Spectrometry

Derivatized sterol samples were analyzed by on Thermo Scientific GCMS system (Trace 1300 ISQ QD). Samples were injected in split mode at 250°C with a ratio of 1:10 and helium at a flow rate of 1.4 mL per min was used as carrier gas. Starting temperature was held at 50°C for 1 min, ramped at a rate of 50°C/min upto 260°C, then again ramped at 2°C/min upto 325°C and held for 3 min. Masses were detected within a range of 50 to 600 atomic mass units using electron impact ionization (70 eV). Transfer line and the detector were operated at 325°C and 250°C respectively.

Different sterol intermediates were identified by their retention times with respect to the internal standard, cholestane and fragmentation patterns using Chromeleon 7 (Thermo Scientific). The spectra obtained were matched to NIST library and those described by Muller *et al* [25]. The relative abundance was calculated from the peak area of each sterol molecule normalized to the internal standard signal.

2.8. Protein Estimation.

An aliquot of 25 μ L was taken from lysate of each replicate for total protein estimation in order to normalize the quantified data. Protein estimation was done using bicinchoninic acid (BCA) protein assay kit (G-Biosciences, MO, USA) as described previously [26].

2.9. Data Analysis.

Mass spectrometric chromatograms of SL and PL molecules were analyzed by MultiQuant™ software (SCIEX). As mentioned earlier, separate internal standards for SLs and PLs were used to normalize the area of each lipid species. Three biological replicates were run for all analyses and data was presented as % of total SL, PL or sterol content. Data was plotted using GraphPad Prism 8. Structures of lipid molecules were drawn using structure drawing tools available on <https://www.lipidmaps.org/resources/tools/structure>.

3. Results

3.1. *R. delemar* Harbours All Major SL Classes

The relevance of SLs in *R. delemar* became apparent when we checked its susceptibility towards specific inhibitors. *R. delemar* cells were susceptible to myriocin, a potent inhibitor of serine palmitoyltransferase (SPT), which is the enzyme responsible for the first and rate-limiting step in SLs biosynthesis (**Figure 1a**). By inhibiting SPT, myriocin disrupts SLs homeostasis in yeast cells, which can have various downstream effects on cell physiology and function [27]. On the other hand, *R. delemar* cells were resistant to aureobasidin A (AbA), a cyclic depsipeptide antibiotic that inhibits inositol phosphorylceramide (IPC) synthase encoded by the *AUR1* gene (**Figure 1a**). The resistance to AbA indicated a lack of acidic SL pathway in *R. delemar* cells (discussed below). Our susceptibility profile of *R. delemar* underscores the relevance of SLs in its various cellular processes which prompted us to analyse SL composition in detail. SLs were detected in base hydrolyzed lipid samples by Shotgun and MRM approaches described previously [22,28]. Multiple confirmatory scans specific for the detection of different SL intermediates were performed in positive mode ($[M+H]^+$). Since it was

the first high throughput analysis of this fungal pathogen, we subjected the samples to unbiased scanning over mass range of m/z 200 to 1000 dalton (Da) which included most of the commonly found SLs in fungi. Separate chromatograms were recorded for each type of scan, which included numerous masses. The mass signals obtained in each chromatogram corresponded to SL molecular species representing the different SL classes which are structurally unique from each other. The different confirmatory scans and the detected SL intermediates are given in **Supplementary File S1**.

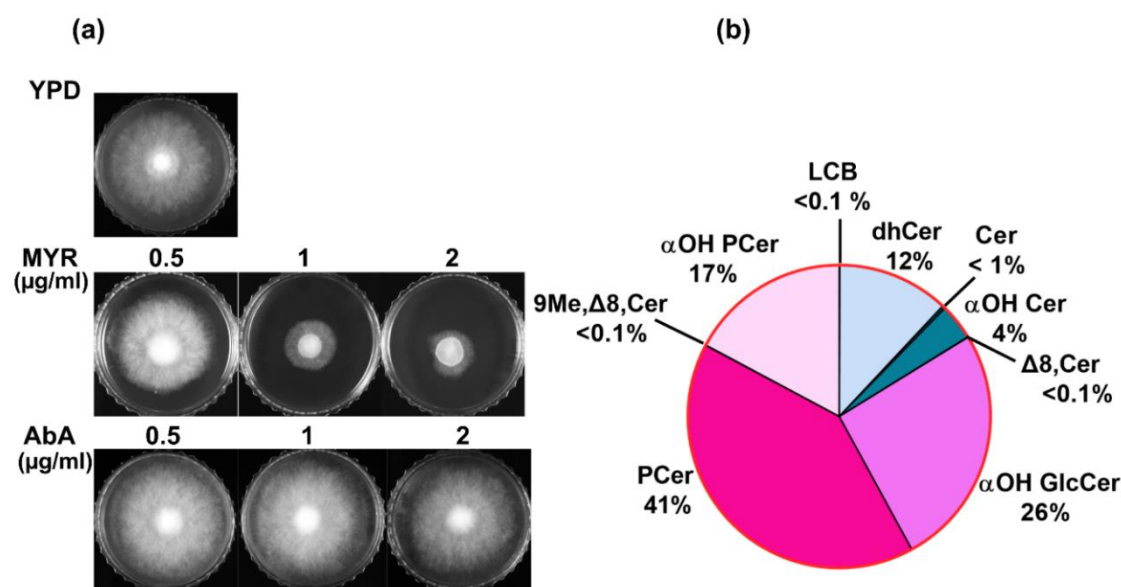


Figure 1. a. Susceptibility of *R. delemar* against SL inhibitors MYR and AbA. Approx. 10^3 spores were spotted on YPD plates with different concentrations of MYR and AbA. Plates were imaged after incubating at 30 °C for 48 hours. **b.** Pie-chart represents proportion of different SL intermediates in *R. delemar*.

Our analyses of the detected masses enabled us to identify all diverse classes of SLs that differed based on headgroup, presence of double bonds and number of hydroxyl groups on the sphingoid base (backbone). These included commonly found SL classes such as long chain bases (LCBs), dihydroceramides(dhCer), ceramides, alpha-hydroxy ceramides(α -OH Cer), alpha-hydroxy glucosylceramides (α -OH GlcCer), phytoceramides (PCer) and alpha-hydroxy phytoceramides (α -OH PCer) [22,26,28]. In addition, fungal specific lipids such as Δ 8-ceramide and 9-methyl, Δ 8-ceramide were also detected (**Figure 1b**). Additionally, all the molecular species detected had five types of backbones. Also the fatty acids acylated to sphingoid base were either saturated, or mono-unsaturated, hydroxylated (likely at C-2 position) or non-hydroxylated thus combinedly generating multiple pools of SL species. Selected masses from the shotgun approach were further detected via MRMs and the samples were subjected to LC-ESI/MSMS for separation and quantification of each lipid species. Peak identification and quantification of each species was done on the basis of retention time (RT) with respect to available natural SL standards. The complete list of identified SL species and their relative abundance is given in **Supplementary File S2**

3.1.1. Long Chain Base (LCBs) Content Is Very Low with DHS-1-P as the Major Base.

Our analysis could detect all major LCBs including dihydrosphingosine (DHS), phytosphingosine (PHS), sphingosine (SPH), dihydrosphingosine-1-phosphate (DHS-1-P), sphingosine-1-phosphate (SPH-1-P). Notably, glucosyl-sphingosine (Glu-SPH) and phytosphingosine-1-phosphate (PHS-1-P) could not be detected in *R. delemar* cells (**Figure 2**). Combinedly, the total LCB content was very low (0.05 %) with DHS-1-P emerging as a major base (0.02%) followed by PHS (0.01%), both sharing approximately two-thirds of the LCB content between them. Other bases such as DHS, SPH and SPH-1-P were present in trace amounts ranging between

0.001% and 0.009%. In comparison with other yeasts such as *Candida* and *Cryptococcus* wherein LCBs ranged between 0.5% and 1.2%, its levels are very low in *R. delemar* cells [22,26,28].

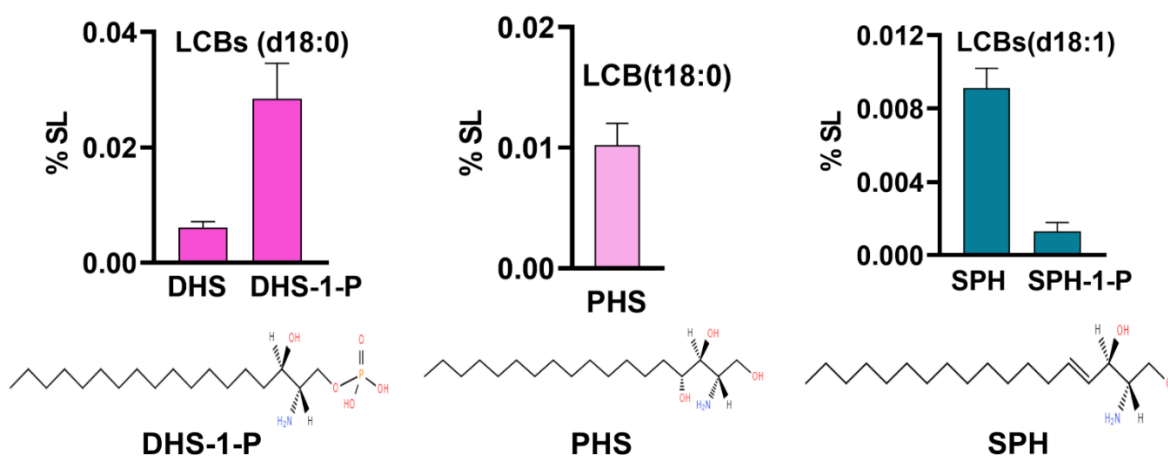


Figure 2. LCBs detected in *R. delemar*: DHS, DHS-1-P(d18:0), PHS(t18:0) and SPH, SPH-1-P(d18:1). Bar-graphs represent relative proportion of these bases out of total SL content. Structures represent top three major bases, each with different backbone.

3.1.2. dhCer Synthesis Leads the Initiation of Neutral Pathway and 16:0 Is the Major Species.

dhCer structures are synthesized by the acylation of the sphingoid base, DHS, through the amine group present at C2-position of the base. These intermediates serve as starting precursors of the neutral branch of the fungal SL biosynthesis pathway. We could detect multiple species of dhCer. For the analysis, selected masses corresponding to molecular species with of 16 to 28 carbon long fatty acid chains were quantified (Figure 3a). Total dhCer content was ~12% with 16:0 dhCer as the predominant species (Figure 3b). Notably, the abundance of 16:0 species is different from other species of fungi such as *Candida*, *Cryptococcus* where 18:0 dhCer predominates[22,28]. The species with longer chain lengths, 24:0, 26:0 and 28:0 shared 3%, 3% and 0.7%, respectively of the SL content. Other saturated fatty acid containing species such as 18:0, 20:0, 22:0 were present in very low amount, each having $\leq 0.1\%$ contribution. Mono-unsaturated fatty acid containing species were present in trace amounts.

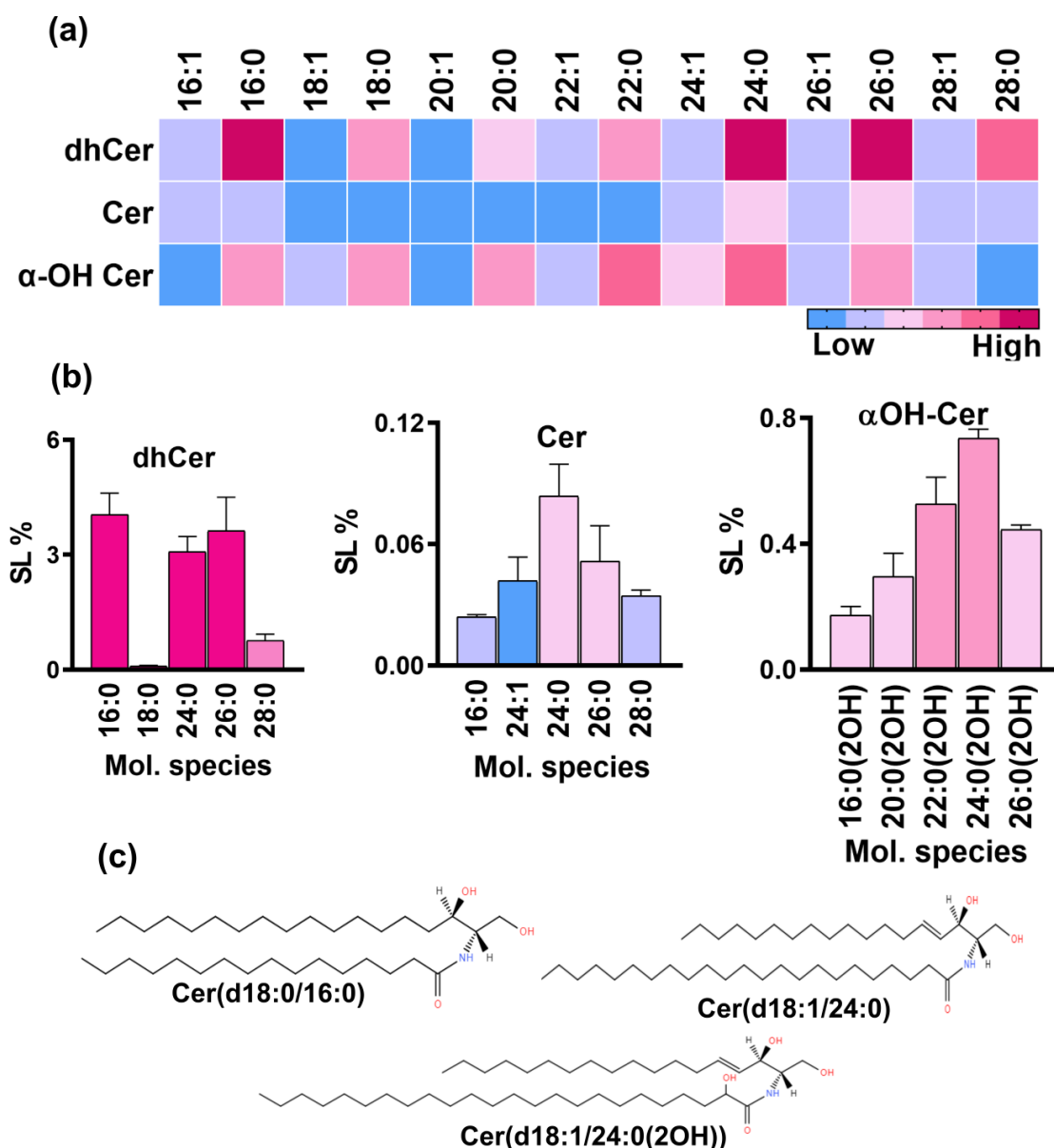


Figure 3. SL classes and molecular species of the neutral branch of SL biosynthesis pathway in *R. delemar*. a. Heatmap represents the relative representation of different molecular species of dhCer, Cer and α OH-Cer. b. Top five species of each class were quantified and represented by bar-graphs. c. Structures represent the major molecular species of each class.

3.1.3. Cer and α -OH Cer Species Were Also Detected and 24:0 Is the Major Specie in Both Classes.

The desaturation of dhCer (d18:0 backbone) leads to the formation of Cer which is a central hub acting as a precursor of many simple to complex SL in eukaryotic cells. The total Cer content was ~0.3% mainly contributed by species with 16C to 28C long fatty acids, out of which 24:0 was the major species (0.08%) followed by 26:0, 24:1, 28:0, 26:1, 16:0 and 16:1 (ranging between 0.02 and 0.05%) (**Figure 3b**). The other Cer species were found in trace amounts. Notably, mono-unsaturated fatty acid containing species were very low than the saturated ones with the same carbon number chain length. SLs with same backbone as that of Cer (d18:1), but having hydroxy-fatty acids most likely at the C2- position, known as α OH-Cer were also present. Total α OH-Cer was ~3.9 % of the total SL content with a wide range of species ranging from 14C to 30C long fatty acids. The major species were 24:0, 22:0, and 20:0 contributing 1.2%, 0.7%, 0.5% and 0.4%, respectively. The other prominent species were 20:0, 18:0, 16:0 and 14:0 ranging from 0.1% to 0.29%. Other species and the mono-unsaturated fatty acid containing species were present in very low amounts (<0.1%).

3.1.4. *R. delemar* Harbours Typical Fungal Specific Lipids

The fungal specific SLs, di-unsaturated sphingoid base, d18:2 or $\Delta 8$ -Cer synthesized by desaturation of Cer at C8-position were detected (**Figure 4**) along with its downstream intermediate, 9-methyl, $\Delta 8$ -Cer having d19:2 backbone [29]. Both types of SLs are unique to certain fungi such as *Candida* and *Cryptococcus* and are not reported in *S. cerevisiae* and mammals [30]. In d18:2, two Cer species with non-hydroxy fatty acids, 16:0 and 16:1 and two species of α -OH-Cer, 16:0(2OH) and 18:0(2OH) were quantifiable. The former two represented an amount of 0.001% to 0.01% while as the latter two were 0.02% and 0.0001%- thus again showing that 16C containing species predominate others as in dhCer. In 9-methyl, $\Delta 8$ -Cer, five species were detected, out of which three were non-hydroxy fatty acid species (d19:2 Cer) and two were α -OH Cer species (d19:2 α -OH Cer). The d19:2 Cer species include 16:0, 16:1 and 18:0 representing 0.004%, 0.01% and 0.001%, respectively. The two α -OH Cer species 16:0(2OH) and 18:0(OH) were 0.02% and 0.0007% respectively. Species with d18:2 and d19:2 backbone are precursors of neutral complex SLs such as GlcCer.

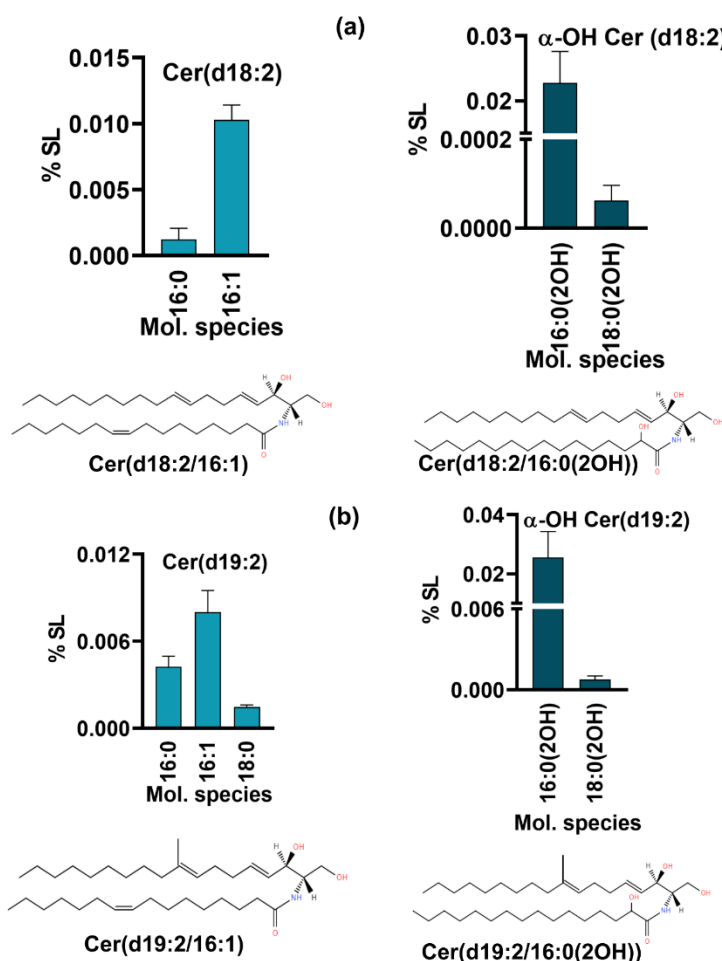


Figure 4. Fungal specific SL intermediates. **a.** having d18:2($\Delta 8$ -Cer) **b.** having d19:2 backbone (9Me, $\Delta 8$ -Cer). Bars depict the top molecular species from each backbone type. The structure of one representative specie is depicted at the bottom of **a** and **b** panels.

3.1.5. Glucosylceramides Represent Second Most Abundant SL Class

As part of the neutral SL biosynthesis pathway, SL intermediates with d18:2 and d19:2 backbones are glycosylated at the C1-position to yield fungal complex SLs such as hexosylceramides with glucose as most common sugar in the headgroup. Our analyses detected α -OH GlcCer as a major complex SL representing ~26% of the total SL content, thus making it second most abundant SL class

in *R. delemar* after PCer. A total of five molecular species were detected that showed significant abundance (**Figure 5**). Two GlcCer species from the d18:2 pool had 16(2OH) and 18:0(2OH) as fatty acid contributing 0.3% and 0.01%, respectively. In contrast, three GlcCer species from the d19:2 pool had cumulatively more abundance than the d18:2 pool and had 16:0(2OH), 18:0(2OH) and 24:0(2OH) as the major species sharing ~25%, 0.1% and 0.01% of the total SL content among them. Infact GlcCer d19:2/16:0(2OH) species was the single largest among all the species detected interms of quantity. This observation differs from other fungi such as *Candida* where 18:0 (2OH) is the major GlcCer species [22,28]. GlcCer synthesis represents the terminal step in the neutral branch of the SL biosynthesis pathway.

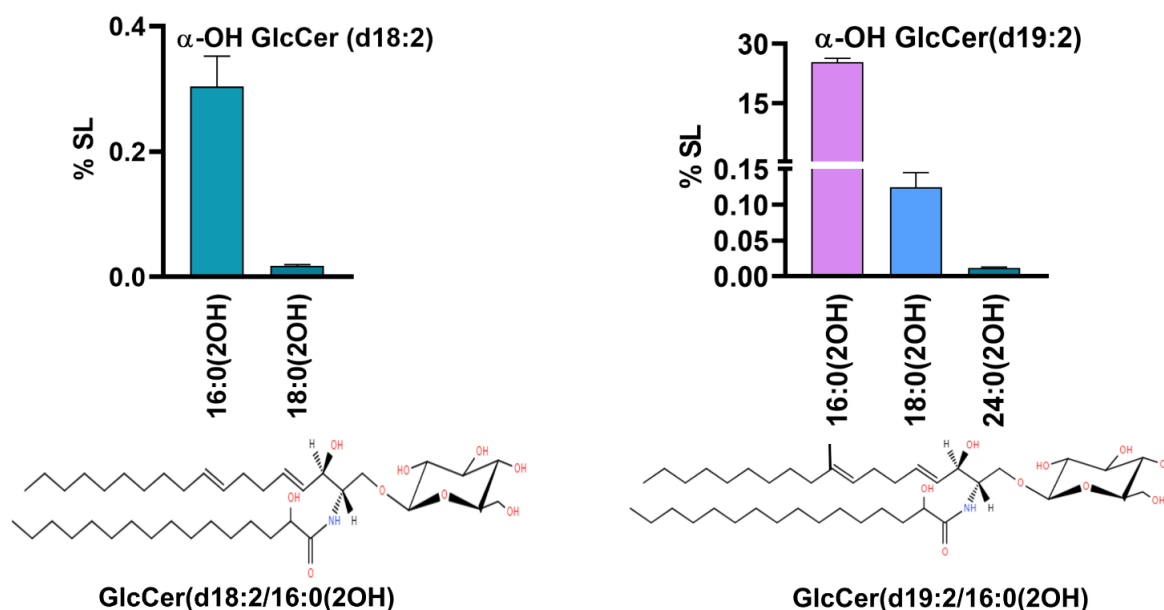


Figure 5. Five major species of α OH-GlcCer with d18:2 and d19:2 backbones acylated to fatty acids of varying chain lengths (x-axis). Structures of two major α -OH GlcCer species are drawn. GlcCer(d18:2/16:0(OH)) on the left has a backbone of 18 Carbons with double bonds at C-4 and C-8 position, glucose at C-1 position and hydroxyl group at C-2 position of the 16C fatty acid chain. Similarly on the right side, GlcCer(d19:2/16:0(OH)) is the major specie with methyl group at C-9 position of the sphingoid base.

3.1.6. PCer Is the Most Abundant SL Class While Its Terminal Acidic Intermediates Are Absent in *R. delemar* Cells

PCer is synthesized by acylation of PHS via amide linkage at C2-position of the PHS base, thus initiating the t18:0 branch of the SL biosynthesis pathway. It is the important intermediate of acidic SLs pathway. Scanning of the t18:0 backbone having m/z 282.2 yielded masses belonging to many SL classes (**Supplementary file S1**). Most mass signals corresponded to species belonging to PCer which upon quantification revealed PCer as the most abundant SL class with a proportion of 41% out of the total SL content. The detected PCer species prominently included even number of carbons containing fatty acids having chain length of 16 to 28 carbons (**Figure 6a**). The significant species were 26:0, 24:0, 28:0, and 16:0, contributing 19.13%, 13.8%, 5.7%, and 1.1%, to the sphingolipid content, respectively (**Figure 6b**). Other detected species such as 18:0, 20:0, 22:0 and mono-unsaturated fatty acid containing PCer species accounted for a very minute amount (<1%). Notably, species with saturated fatty acids were remarkably higher than the ones having unsaturated fatty acids.

Table 2. position of the fatty acids yields α OH-PCer which serve as precursor of inositol phosphorylceramides (IPCs) intermediates and represents the acidic branch of SLs biosynthesis in yeast and some other filamentous fungi. The acidic SLs are commonly found in several *Candida* and in other fungal species; however, IPCs were not detected in *R. delemar*. The analysis showed the

accumulation of α OH-PCer which was ~17% of the total SLs and thus making it third most abundant SL class (**Figure 1b**). Multiple molecular species of α OH-PCer with fatty acids of varying chain length (14C to 28C) were detected and quantified (**Figure 6a**). Like PCer, major specie had similar number of carbons in fatty acids but hydroxylated such as 24:0(2OH), 26:0(2OH) representing 9.5% and ~7% of the total SL content, respectively (**Figure 6b**). These were followed by 24:1(2OH), 26:1(2OH) and 28:0(2OH) with amount of 0.16%, 0.13% and 0.02%, respectively. Other species such as 14:0(2OH), 16:0(2OH), 18:0(2OH), 20:0(2OH), 22:0(2OH) and other mono-unsaturated species were only found in trace amounts ($\leq 0.01\%$).

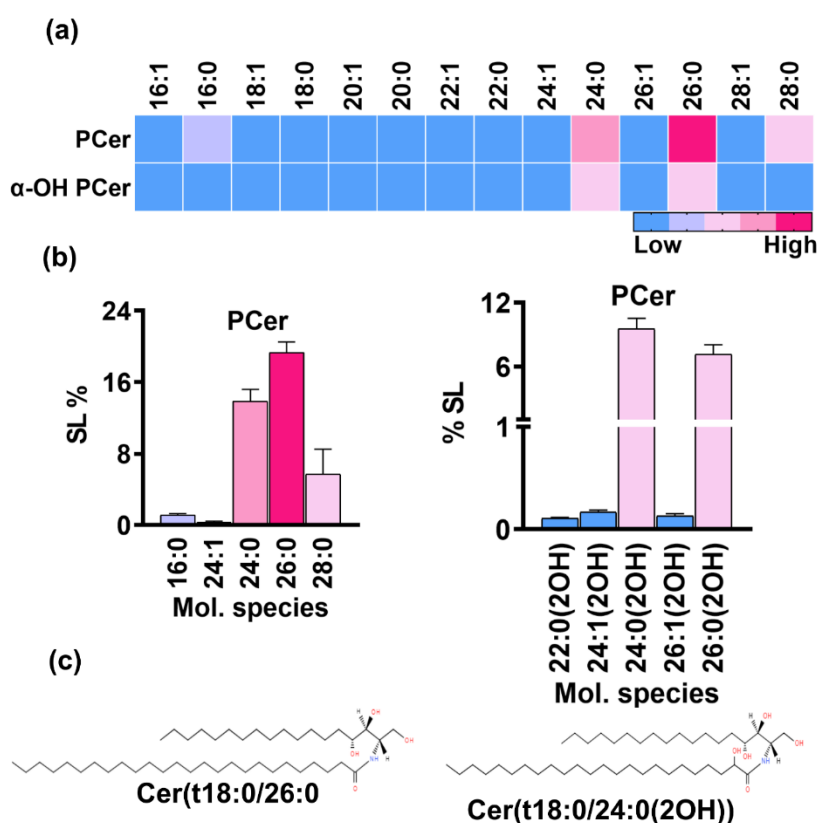


Figure 6. a. PCer and α OH-PCer, two major classes in the acidic branch and their molecular species. **b.** Bargraphs represents the relative amounts of top five molecular species from each class. At the bottom, structures of major species in each class are shown.

3.2. All Major Glycerophospholipids (GPL) Are Present in *R. delemar*.

Considering that GPL play diverse and important roles in the biology and pathogenesis of Mucorales, we analysed it in great detail wherein total extracted lipids were subjected to shotgun lipidomics in which different confirmatory scans targeting the polar headgroups of GPLs were used for detection of particular classes [31]. All scans were performed in positive mode ($[M+H]^+$ or $[M+NH_4]^+$) and ranged between m/z 400 Da to 1000 Da covering all the known PL species. The different GPLs detected are given in **Figure 7**.

Multiple species were detected that belonged to six classes of phospholipids (PLs). Approximately 200 top species were selected and quantified using MRM approach in which separate chromatograms were recorded for each species and the identification was done on the basis of retention time with respect to the available natural standards of each class. Each class has a unique polar headgroup and one or two acyl chains linked to glycerol backbone via ester linkages. The complete list of quantified PL species is given in **Supplementary file S2**.

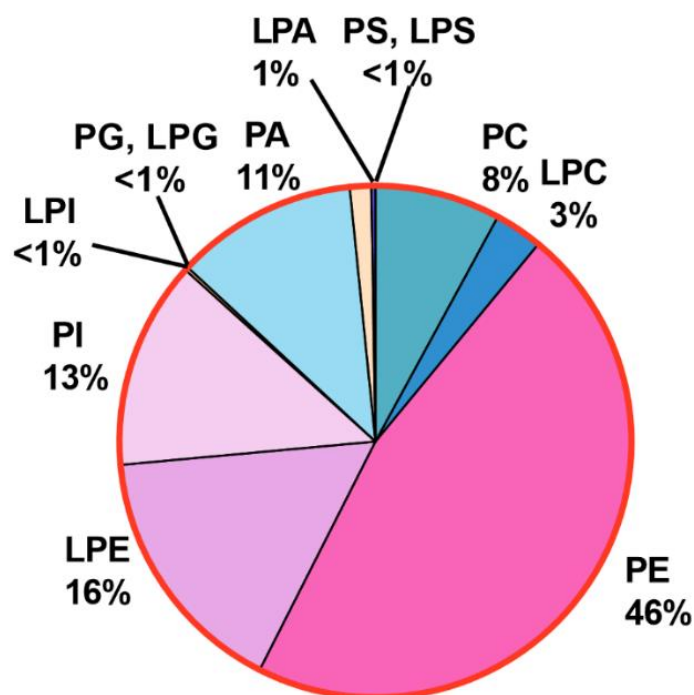


Figure 7. PL landscape of *R. delemar* with relative distribution of PL classes.

3.2.1. Phosphatidylethanolamine (PE) and Lysophosphatidylethanolamine (LPE) Constitute the Bulk of PL Content

PL classes with ethanolamine headgroup and having two acyl chains i.e; PE was found to be the most abundant PL class (46%). Together with LPE (16%) having only one acyl chain, these combinedly contribute bulk of the total PL content (**Figure 8**). Both types of species were detected using a Neutral Loss Scan of 141 Da. While our analysis could detect numerous PE species, those with 30 to 38 carbons and having total double bonds ranging from 0 to 6 in both the acyl chains showed significant abundance and were quantified (**Figure 8a**). Notably, species with unsaturated fatty acids were more abundant than those with saturated fatty acids. For instance, the major PE species found had combinedly 34C in the acyl chains such as PE 34:1(11.6%), 34:2(11.2%), 34:3(5.1%) (**Figure 8b**). This was followed by species with 36C such as PE 36:2(4.4%), 36:4(3.4%), 36:3(3.2%), 36:1(~2%) and 36:5(1.8%) and 32:1(1.8%). All other species were <1%. Species with odd-chain fatty acids with total 33C and 35C were also detected and were present in low amounts.

LPE species were also present in high quantities (**Figure 8a**). Five major species were detected, out of which 18C containing species predominated. LPE 18:1 was most abundant (7.3%) followed by 18:2 (5.4%) and 18:3 (1.36%) (**Figure 8b**). Similarly 16C containing LPE 16:1 was the main species followed by 16:0 sharing ~1% and 0.4% PL content, respectively.

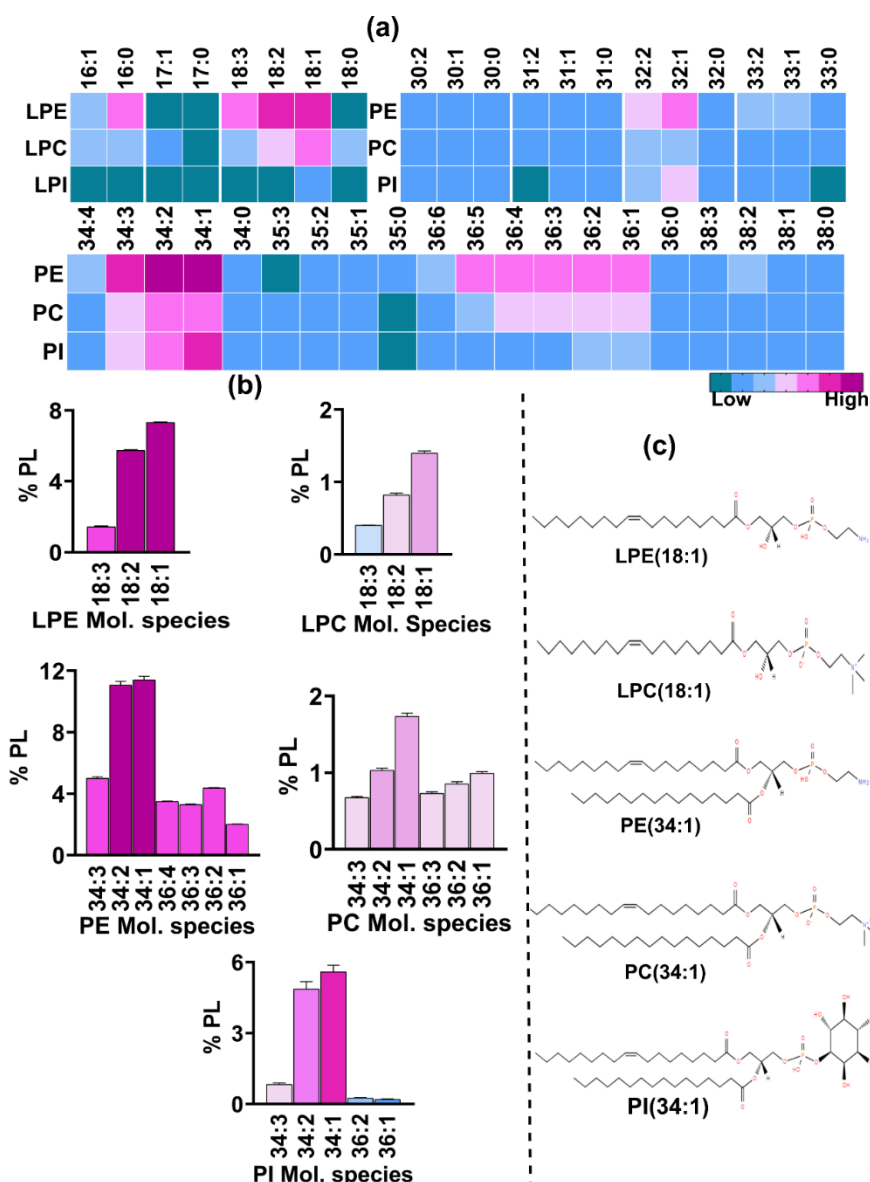


Figure 8. Major PL and Lyso-lipid species of different PL classes detected in *R. delemar*. **a.** Heatmap represents relative abundance of molecular species in each class with varying number of carbons and double bonds in both acyl chains. **b.** Top molecular species from PC, PE, PI were quantified and represented by bar-graphs. **c.** Structures represent major molecular specie from each class.

3.2.2. PhosphatidylCholine (PC) Content Is Low

Unlike many fungal species, PC contents are low in *R. delemar* cells. The precursor ion scan of phosphocholine headgroup (m/z 184.1 Da) revealed presence of species belonging to PC and LPC. Cumulatively, both types contributed ~11% to the PL content out of which PC was ~8% and LPC 3%. Like PE, species contained a total of 30C to 38C out of which 34C containing species, such as PC 34:1, 34:2 were most abundant, having a proportion of 1.7% and 1%, respectively (**Figure 8b**). This was followed by species with 36C such as 36:1(~1%), 36:2(0.85%) and 36:3(0.73%). Two other significant species included 34:3(0.65%), 36:4(0.56%), 32:1 and 36:5 each sharing 0.3% of the total PL content. Other species including those with saturated fatty acids and with odd chain fatty acids of 31C, 33C and 35C were very low(≤ 0.1).

Similarly in LPC, species with saturated and unsaturated fatty acid chain length of 16C to 18C were detected in which LPC 18:1, 18:2 and 18:3 were major species with 1.8%, 0.8% and 0.4 %

proportion, respectively, followed by LPC 16:0, 18:0 and 16:1 ranging from 0.12 % to 0.16% (**Figure 8b**). Odd chain containing LPCs, 17:0 and 17:1 were found in trace amounts.

3.2.3. Phosphatidyl Inositol (PI) and Phosphatidic Acid (PA) Show Equal Abundance

Lipid species having inositol as a headgroup were detected using the confirmatory Neutral Loss scan of 277 Da in positive mode. Multiple species belonging to PI having two acyl chains were detected and quantified (**Figure 8a**). However, lysophosphatidyl Inositol (LPI) which has only one acyl chain was scarce as only one LPI species showed significant presence. All PI and LPI species constituted about 13% of the total PL content. The major species among PI were those having a total of 34 carbons in both acyl chains, such as PI 34:1, 34:2 and 34:3 contributing 5.5%, 4.8% and 0.84%, respectively, to the total PL content (**Figure 8b**). These were followed by 32C containing PI 32:1 and 32:2, having a proportion of 0.5% and 0.4%, respectively. Unlike PC and PE, 36C containing species were less abundant than those with 32C. Major species in 36C group were having one to five double bonds with their levels ranging from 0.05% to 0.2%. Other species including the odd chain fatty acid containing PI species and a single LPI species, LPI 18:1 were present in trace amounts.

Similarly lipid species with phosphate headgroup were detected by Neutral Loss of 115 Da scan and included PA and lyso-PA (**Figure 9a**). Their combined abundance of 12.8% was nearly equal to PI and LPI (~13%). The major PA species were PA 34:2, 34:1 and 34:3 having proportion of ~2.4%, 2.2% and 0.7% respectively (**Figure 9b**). These were followed by species having 36C and one to five double bonds with their proportions ranging from 0.5% to 1.2%. Species from the 32C group, PA 32:1 and 32:2 were 0.45% and 0.3%, respectively. All other PA species were present in low amount ($\leq 0.1\%$).

Unlike LPI, five species of LPA had significant presence with major species having 18C and 16C in their acyl chain. The major species were LPA 18:1(0.7%),18:2 (0.5%), 16:1(0.04%),16:0 (0.02%) and 18:0 (0.01%).

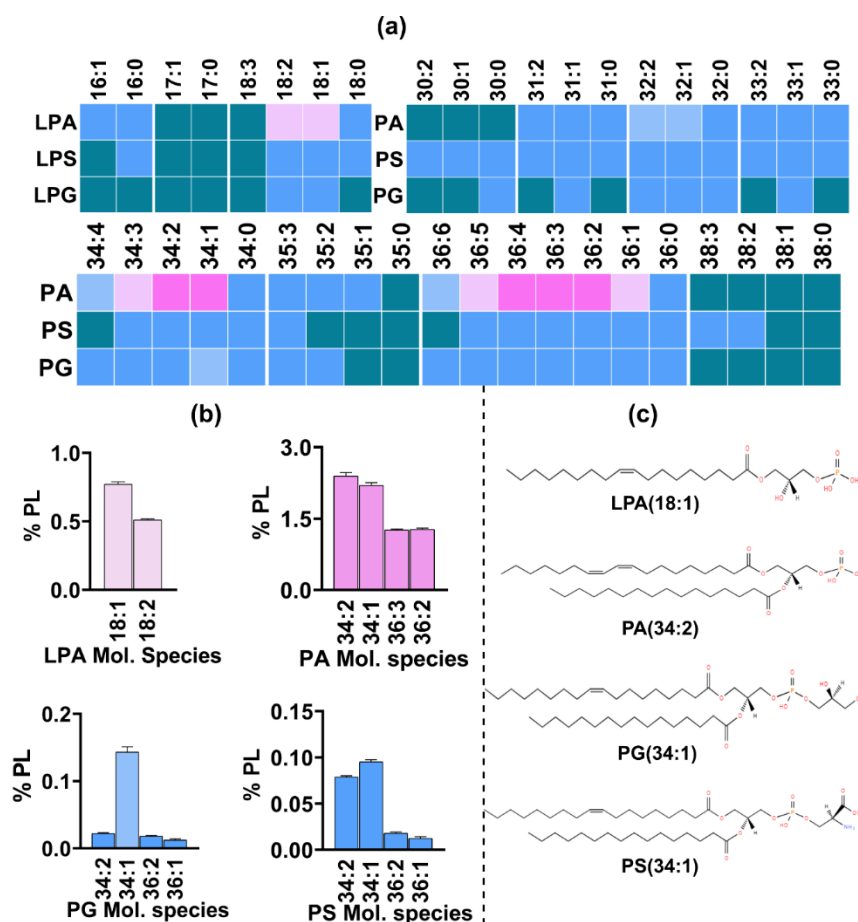


Figure 9. Major PL classes detected in *R. delemar* and their Lyso-species. **a.** Heatmap represents relative abundance of molecular species in each class with varying number of carbons and double bonds in both acyl chains. **b.** Abundant molecular species from PA, PG, PS were quantified and represented by bar-graphs. **c.** Structures represent major molecular species from each class.

3.2.4. Phosphatidyl Serine (PS) and Phosphatidyl Glycerol (PG) Represent the Least Abundant GPLs.

In the positive mode, a neutral loss scan of 185 Da detected lipid species with serine headgroup.. Species belonging to PS with a total of 30C to 38C in their acyl chains showed significant presence (**Figure 9a**). The pattern was similar as that of other PL classes with 34C containing species being more abundant than 36C and 32C containing species (**Figure 9b**). The major species from the 34C group were PS 34:1, 34:2 and 34:3 having a proportion of 0.1%, 0.08% and 0.02%, respectively, followed by species within 36C group such as PS 36:2, 36:1, 36:3 with their respective amounts as 0.02%, 0.01% and 0.01%. Among the 32C group, PS 32:1 showed significant presence (0.01%). All other PS species including saturated and odd chain fatty acids were present in trace amounts (<0.01%). Four LPS species LPS 16:0, 18:0,18:1, and 18:2 were also found in minute amounts. PS and LPS combinedly constituted about 0.2% of the total PL content.

Species with glycerol as headgroup were also detected by Neutral loss of 189 Da scan, and belonged to either phosphatidyl glycerol (PG) with two acyl chains or lyso PG having only one acyl chain. Similar to PS and LPS, the total share of PG and LPG was found to be ~0.2%, hence making it the least abundant PL class. The compositional distribution of PG species also followed the same pattern as that of other PL classes with PG 34:1 and 34:2 being most abundant with their proportion as 0.1% and 0.02%, respectively (**Figure 9b**). These were followed by PG 36:4 and 32:1 sharing 0.006% and 0.005%, respectively. All other species showed very low presence. Among LPG, two species, LPG 18:1 and LPG 18:2 were quantifiable and ranged between 0.001% to 0.008%.

3.3. Ergosterol Is the Major Sterol and Alternative Sterol Biosynthesis Pathway Also Exists in *R. delemar*

We employed GCMS to detect and identify major intermediates in the sterol biosynthesis pathway in *R. delemar*. Sterols were identified as TMS-derivatives and spectral pattern of each peak was searched against the spectral libraries of sterols [25]. While we could detect a number of TMS-derivatized sterols, some non-derivatized and unidentified sterols were also detected. Here, we selected and quantified top sterol intermediates in which ergosterol was found to be the most abundant sterol (~90%), an observation widely found in fungal kingdom (**Figure 10**). This was followed by ergosta-5,7 dienol (~7%) and its upstream intermediate, episterol (~1%). The first sterol in the fungal biosynthesis pathway sequence lanosterol was also detected. Its downstream products eburicol and zymosterol were also detected and also showed significant abundance. Both intermediates via fecosterol lead to synthesis of ergosterol through multiple steps.

Notably, cholesta type sterols synthesized from zymosterol were also detected. Major cholesta type sterols were cholesta-7,24 dienol, cholesta-5,7,24 trienol and cholesta-5, 7, 22, 24 tetraenol. These intermediates are part of the alternative pathway of ergosterol biosynthesis in *R. delemar* (**Figure 10**).

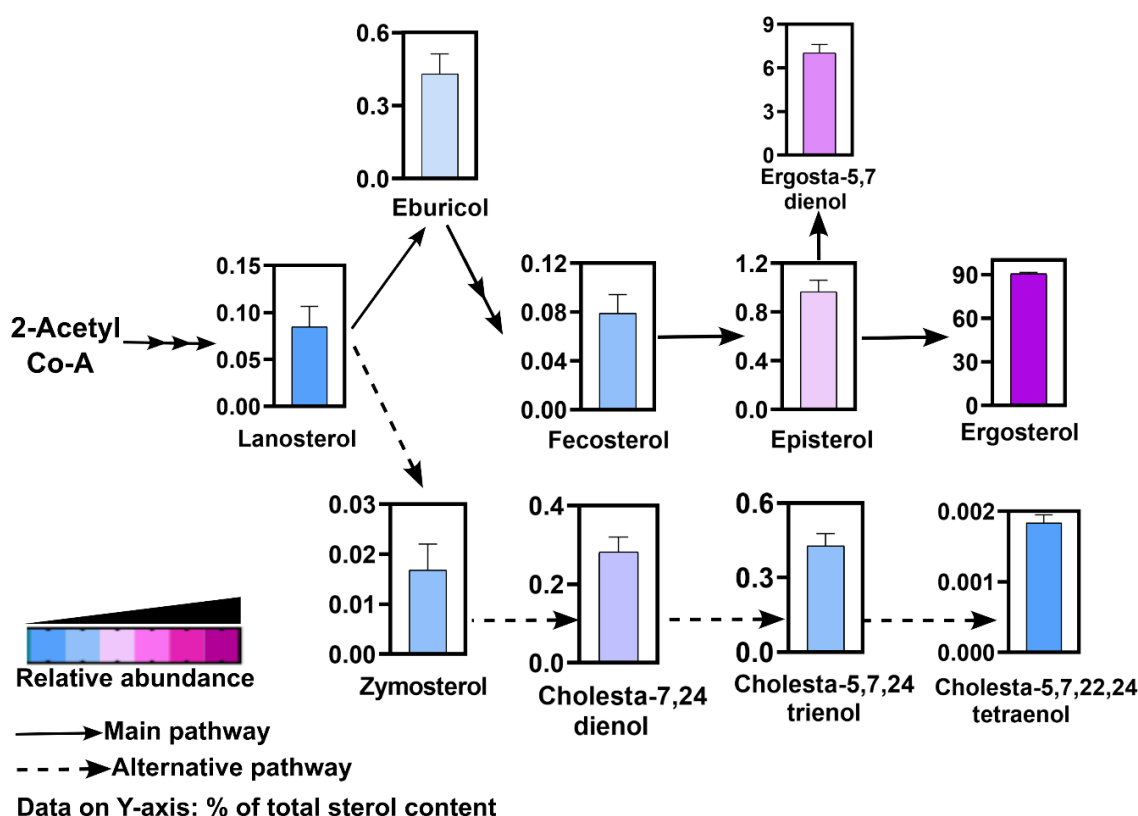


Figure 10. Major sterol intermediates in *R. delemar* as detected by GCMS. Only the detected intermediates have been shown as bargraphs. Data on Y-axis represents % of each sterol intermediate of total sterol content.

Discussion

Membrane lipids are targets of the common antifungal agents such as azoles and polyenes. Previous studies from others and our group suggest that perturbations in lipid profiles of fungi leads to antifungal resistance development [32–36]. Considering the intricate relationship between lipids and drug resistance, *R. delemar* 99-880, a genome sequenced [21,37,38] and widely characterized representative of Mucorales was used to study compositional landscape of lipids. For the first time, we dissected three major lipid biosynthesis pathways such as SLs, PLs and sterols and not only detected major intermediates of their respective pathways, but also quantified major classes and their constituent molecular species that differed in carbon chain length, and enzymatic modifications such as hydroxylations and degree of unsaturation. The predicted biosynthetic pathways in **Figure 11**, are based on detection of the lipid intermediates and their relative abundance.

Our analysis confirms an earlier study by Aoki et al., which demonstrated evidence of AbA resistance in many Mucorales species, as well as in ascomycete fungus *Hirsutella rhossiliensis* and is due to absence of typical acidic SLs such as IPCs and mannosyl inositol phosphorylceramides (MIPCs) [39,40]. Also neutral SLs species having one to many sugar molecules in GlcCer have been described in the same study [39]. However, information about their upstream precursors remained unexplored. In our analysis, we confirmed the presence of fungal specific SLs of the neutral pathway with d18:2 and d19:2 sphingoid backbones, that are precursors of complex GlcCers (**Figure 11a**). GlcCer probably being the sole complex SL in *Rhizopus* could play critical roles. In *Cryptococcus*, GlcCer has already been shown to be a major virulence factor as its depletion rendered cells avirulent [41]. In other filamentous fungi, GlcCer is required for polarized growth, spore germination and alkali tolerance while in *C. albicans*, GlcCer mediates hyphal growth and also susceptibility towards SDS, and FLC [30,42–44]. Our recent studies have also suggested GlcCer compositional differences associated with drug resistance in clinical isolates of *C. auris* [26]. The lower abundance

of upstream precursors such as d18:2 and d19:2 Cers and OH-Cer reinforces that these intermediates are highly fluxed towards synthesis of GlcCer to maintain its required proportion. Notably, exogenous GlcCer has been shown to antagonize the effects of the potential antifungal drug, miltefosine, in many species of Mucorales [45]. Similarly, among LCBs, DHS-1-P was found to be major base compared to PHS and DHS which are found to be abundant in *Candida* [26,46,47]. *LCB4*, encoding gene for LCB kinase is part of *Tac1* regulon, which is one of the major determinants of azole resistance in *Candida* [48]. Phosphorylated bases are known to be involved in temperature tolerance in *S. cerevisiae*, although their roles in other fungi are still unclear [49].

Polar lipids such as PLs detected in our analysis were structurally similar to other eukaryotes. However, remarkable differences with regard to their relative composition among different PLs classes exist. For instance, the major characteristics observed in *R. delemar* PLs is a high proportion of PE as compared to lower levels of PC, and high abundance of lyso-lipids (**Figure 11 b**). Such apparent variations reflect a completely different physiological state of *R. delemar* compared to other fungal pathogens, where PC usually predominates among all other lipid classes [50]. Notably, a high PE content was seen to be associated with hyper-virulence in *C. albicans* [51]. In addition, a high PE:PC ratio is a major determinant of membrane rigidity that governs drug diffusion across the membrane and affects susceptibility [36,52]. Certain lyso-phospholipids were observed to be highly enriched in drug resistant *C. auris* isolate [46]. The high abundance of lyso-lipids such as LPE in *R. delemar* suggests strong phospholipase activity which impact fungal fitness and virulence. Lyso-lipids were also found in *Candida* extracellular vesicles in response to oxidative stress [53]. PLs are known to play roles in maintain membrane rigidity, cell cycle progression and also govern stress resistance. In one of our previous studies, drug resistant *C. auris* isolate had higher PL levels compared to susceptible isolate [54].

Sterol metabolism in fungi and its role in antifungal drug resistance has been a subject of research for a long time. As common antifungal drugs like azoles and polyenes target sterols, any alteration in sterol profile impacts drug susceptibility. Mutation in sterol biosynthesis genes such as *ERG6*, *HMG1* and *ERG3* in *Candida*, *Aspergillus*, *Mucor lusitanicus* and *M. circinelloides* impact susceptibility of azoles and polyenes [55–57]. *Erg6* deletion leads to the synthesis of cholesta-type lipids from zymosterol via an alternative ergosterol biosynthesis pathway-often associated with antifungal drug resistance. Genomic analysis of *R. oryzae* (*R. delemar*) [21] also reveals presence of genes responsible for biosynthesis of alternative sterol intermediates, some of which were confirmed in our study as discussed earlier. Our analysis of measured sterol intermediates confirms all reported alternative sterol pathways which also appear to operate in *R. delemar* (**Figure 11c**). Also, *R. delemar* genome contains multiple copies of several ergosterol biosynthesis genes such as *ERG11*, *ERG6*, *ERG3* and gene duplication of sterol biosynthesis genes has been linked to azole resistance [21]. All these observations point towards potential resistance evolution in *Rhizopus* clinical isolates.

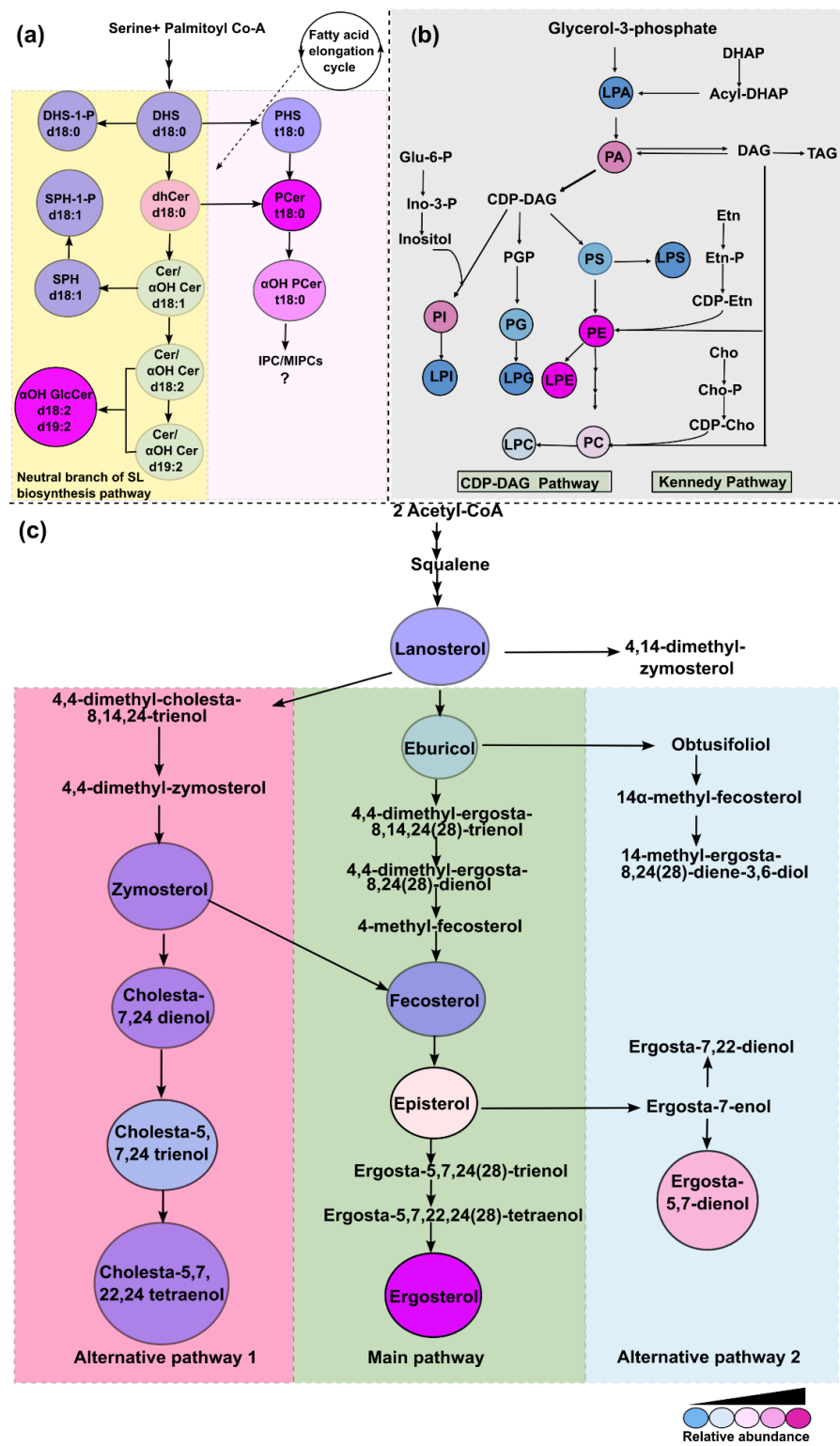


Figure 11. Putative lipid biosynthesis pathways in *R. delemar*. Relative abundance of each detected intermediate is depicted by different colours. **a.** SL biosynthesis pathway based on the intermediates detected in our analyses. Description of SL class in circles is given by short abbreviations followed by characteristic backbone present. **b.** PL biosynthesis pathway and the major classes are given in circles. **c.** Sterol biosynthesis pathway as described in other fungi. Sterol intermediates in the circles are the abundant ones detected in our analysis. The intermediates in the central panel form the main pathway while as the right and the left are alternative pathways.

In conclusion, our study presents the first draft landscape of all major classes of lipids in *R. delemar*, a reference species in Mucorales studies. Analyzing lipid composition among different Mucorales species, which exhibit varying levels of drug resistance, will be interesting since it could explain their diverse physiological roles and reveal their relevance in drug resistance. Nonetheless, the data provided here is expected to serve as a platform to stimulate further research on lipids, their physiological relevance, and their role in drug resistance.

Supplementary Materials: The following supporting information can be downloaded at the website of this paper posted on Preprints.org.

Author Contributions: Conceptualization, B.A., A.S. and R.P.; methodology, B.A., A.C., R.V. and A.S.; software, B.A. and R.V.; validation, B.A., M.K., R.V., A.S., and R.P.; formal analysis, B.A., P.K., H.C., C.L.R., S.M.R., and R.P.; investigation, B.A., A.S., A.S.I. and R.P.; resources, S.M.R., A.S.I., and R.P.; data curation, B.A., A.C., and M.K.; writing—original draft preparation, B.A.; writing—review and editing, M.K., A.S., A.S.I. and R.P.; visualization, B.A., A.S.I., and R.P.; supervision, A.S., S.M.R., P.V.D., A.S.I., and R.P.; project administration, R.P.; funding acquisition, A.S., S.M.R., and R.P. All authors have read and agreed to the published version of the manuscript.

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