



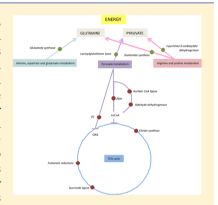
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# Proteomic Profile of Cryptococcus neoformans Biofilm Reveals Changes in Metabolic Processes

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Supporting Information

ABSTRACT: Cryptococcus neoformans, a pathogenic yeast, causes meningoencephalitis, especially in immunocompromised patients, leading in some cases to death. Microbes in biofilms can cause persistent infections, which are harder to treat. Cryptococcal biofilms are becoming common due to the growing use of brain valves and other medical devices. Using shotgun proteomics we determine the differences in protein abundance between biofilm and planktonic cells. Applying bioinformatic tools, we also evaluated the metabolic pathways involved in biofilm maintenance and protein interactions. Our proteomic data suggest general changes in metabolism, protein turnover, and global stress responses. Biofilm cells show an increase in proteins related to oxidationreduction, proteolysis, and response to stress and a reduction in proteins related to metabolic process, transport, and translation. An increase in pyruvate-utilizing enzymes was detected, suggesting a shift from the TCA cycle to fermentation-derived energy acquisition. Additionally, we assign putative roles to 33 proteins previously categorized as hypothetical. Many changes in metabolic enzymes were identified in studies of bacterial biofilm, potentially revealing a conserved strategy in biofilm lifestyle.



KEYWORDS: Cryptococcus neoformans, biofilm, shotgun proteomics, metabolic changes, resistance

# INTRODUCTION

Cryptococcosis is one of the most important systemic mycoses and is considered to be one of the three common opportunistic infections in HIV/AIDS patients, causing almost 625 000 deaths per year worldwide. The importance of cryptococcosis as an opportunistic pathogenic fungus has increased considerably over the last decades due to the intensive chemotherapy of cancer patients, the use of immunosuppressive drugs in organ transplant recipients, and the spread of the AIDS epidemic.<sup>2</sup> This potentially fatal fungal disease is caused by two species of the same genus: Cryptococcus neoformans and Cryptococcus gattii.<sup>3,4</sup>

C. neoformans is an encapsulated opportunistic yeast-like fungus that causes meningoencephalitis in mammalian hosts, mainly in immunocompromised and sometimes in healthy individuals.<sup>5–7</sup> Infection in humans and other susceptible animals occurs by inhalation, establishing a lung infection. To cause central nervous system (CNS) mycosis, C. neoformans becomes blood borne and progresses through until it culminates in fungal replication in the brain. Critical steps include fungal arrest in the vasculature of the brain and interaction and signaling of the fungal and endothelial cells, leading to transmigration with subsequent parenchymal invasion and fungal replication in the CNS.8-10

An ecological strategy that has been associated with chronic infections for several microorganisms, including *Cryptococcus*, is the formation of biofilm.  $^{6,11-13}$  It has been estimated that 65% of all human infectious diseases are biofilm-related. 14 C. neoformans can form biofilms on medical devices, including ventriculoatrial shunt catheters, peritoneal dialysis fistula, cardiac valves, and prosthetic joints. 15-18 Microorganisms growing in biofilms exhibit high resistance to the host immune response, environmental stresses, and antimicrobial therapy. These persistent populations can serve as a reservoir for chronic and systemic infections, playing an important role in human disease. 19,20 This increased resistance suggests that cells in biofilms can modulate metabolic activity, dormancy, and stress responses, 21,22 which highlights the importance of understanding the biofilm-forming properties of *C. neoformans*.

Comparative analysis of the proteomes of several bacterial and fungal pathogens between biofilm and planktonic growth modes has been reported.<sup>23–28</sup> By studying biofilm formation in C. neoformans, we hope to identify general features and specific characteristics of biofilms.

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With the increasing recognition of the role that *C. neoformans* biofilms play during infection, strategies such as functional genomics, proteomics, and metabolomics can help us to gain insight into resistance, antifungal drug targets, and host—pathogen interaction. However, information regarding the molecular mechanisms specifically associated with the biofilm formation remains limited. These approaches could provide a framework for the identification of new proteins and pathways associated with fungal pathogenesis and maintenance.

Here we report the use of shotgun proteomics for comparative analysis of protein expression obtained from biofilm and planktonic cells of *C. neoformans* H99. Also, an interactome analysis revealed differences in protein networks. The changes in protein expression in the biofilm revealed important insights related to energy acquisition, under an oxygen-limiting condition, as indicated by the metabolic pathways analyses, linking the resistance to a persistent infective behavior, a feature also seen in bacterial biofilms.

#### MATERIAL AND METHODS

# **Fungal Strain and Growth Conditions**

Cryptococcus neoformans var. grubii, strain H99 (serotype A) was kindly provided by Dr. Marilene H. Vainstein (Biotechnology Center, Federal University of Rio Grande do Sul, RS, Brazil). The yeast was grown in 50 mL of yeast peptone dextrose (YPD) broth (yeast extract 1%, peptone 1%, glucose 2%) for 20 h, 180 rpm. Cells were separated by centrifugation and washed twice with phosphate-saline buffer (PBS) (10 mM phosphate buffer, 2.7 mM potassium chloride, 137 mM sodium chloride, pH 7.4). The cells were suspended in minimum medium (20 mg/mL thiamine, 30 mM glucose, 26 mM glycine, 20 mM MgSO<sub>4</sub>·0.7H<sub>2</sub>O, 58.8 mM KH<sub>2</sub>PO<sub>4</sub>)<sup>6</sup> and adjusted to the desired cellular density (1 × 10<sup>7</sup> cells/mL) by counting in a Neubauer chamber. This standardized suspension was used as the inoculum to biofilm and planktonic cultures.

C. neoformans biofilms were cultured on Petri dishes containing 20 mL of minimum medium. The plates were maintained in an incubator chamber without shaking for 48 h at 37 °C. After this time, the plates were washed with PBS to remove unattached cells. Then, the resulting biofilm was scraped off the Petri dishes with PBS, transferred to tubes and centrifuged (15 min, 13 000 rpm). For planktonic culture, cells were grown for 48 h at 37 °C with shaking (180 rpm) in 50 mL of minimum medium, and the cells were separated by centrifugation. Both cell pellets were washed twice with PBS, frozen at -80 °C, and lyophilized.

# **Preparation of Protein Extracts**

The lyophilized cells were disrupted with a mortar and pestle in liquid nitrogen to a fine powder, and the samples were suspended in buffer with protease inhibitors (50 mM tris-HCl pH 7.5, 1 mM EDTA, 1 mM PMSF, 50  $\mu$ M TPCK, 5 mM iodoacetamide). Proteins were solubilized by vortexing five times for 1 min each at intervals of 1 min on ice and centrifuged (13 000 rpm for 20 min). Supernatants were collected, and the remaining cell debris were suspended in the same buffer, followed by the same protocol. Supernatants collected after centrifugation were pooled and stored at -80 °C. Protein concentration was determined using the BCA assay (Thermo Scientific, Rockford, IL).

# Sample Preparation for Mass Spectrometry

C. neoformans H99 biofilm and planktonic protein extracts (100  $\mu$ g) were suspended in digestion buffer (8 M urea, 100 mM tris-HCl pH 8.5). Proteins were reduced with 5 mM tris-2-carboxyethyl-phosphine (TCEP) at room temperature for 20 min and alkylated with 10 mM iodoacetamide at room temperature in the dark for 15 min. After the addition of 1 mM CaCl<sub>2</sub> (final concentration), the proteins were digested with 2  $\mu$ g of trypsin (Promega, Madison, WI) by incubation at 37 °C during 16 h. Proteolysis was stopped by adding formic acid to a final concentration of 5%. Samples were centrifuged at 14 000 rpm for 20 min, and the supernatant was collected and stored at -80 °C.

#### MudPIT

The protein digest was pressure-loaded into a 250  $\mu m$  i.d. capillary packed with 2.5 cm of 5  $\mu$ m Luna strong cation exchanger (SCX) (Whatman, USA), followed by 2 cm of 3  $\mu$ m Aqua C18 reversed phase (RP) (Phenomenex, USA) with a 1  $\mu m$  frit. The column was washed with buffer containing 95% water, 5% acetonitrile, and 0.1% formic acid. After washing, a 100  $\mu m$  i.d. capillary with a 5  $\mu m$  pulled tip packed with 11 cm of 3 µm Aqua C18 resin (Phenomenex, USA) was attached via a union. The entire split-column was placed in line with an Agilent 1100 quaternary HPLC and analyzed using a modified 12-step separation as described previously.<sup>31</sup> The buffer solutions used were 5% acetonitrile/0.1% formic acid (Buffer A), 80% acetonitrile/0.1% formic acid (Buffer B), and 500 mM ammonium acetate, 5% acetonitrile, and 0.1% formic acid (Buffer C). Step 1 consisted of a 70 min gradient from 0–100% (v/v) buffer B. Steps 2-10 had a similar profile with the following changes: 3 min in 100% (v/v) buffer A, 3 min in X%(v/v) buffer C, 4 min gradient from 0 to 10% (v/v) buffer B, and 101 min gradient from 10-100% (v/v) buffer B. The 3 min buffer C percentages (X) were 10, 20, 30, 40, 50, 60, 70, 80, 90, and 100% (v/v). An additional step containing 3 min in 100% (v/v) buffer A, 3 min in 90% (v/v) buffer C and 10% (v/v)buffer B, and 110 min gradient from 10-100% (v/v) buffer B were used. Three pools of six biological replicates and two technical replicates were analyzed for both C. neoformans culture conditions (biofilm and planktonic).

# **Mass Spectrometry**

Peptides eluted from the microcapillary column were electrosprayed directly into an LTQ-XL mass spectrometer (Thermo Fisher, USA) with the application of a distal 2.4 kV spray voltage. A cycle of one full-scan mass spectrum (300–2000 *m/z*) followed by five data-dependent MS/MS spectra at a 35% normalized collision energy was repeated continuously throughout each step of the multidimensional separation. To prevent repetitive analysis, dynamic exclusion was enabled with a repeat count of 1, a repeat duration of 30 s, and an exclusion list size of 200. Application of mass spectrometer scan functions and HPLC solvent gradients were controlled by the Xcalibur data system (Thermo, USA).

# **Analysis of Tandem Mass Spectra**

MS/MS spectra were analyzed using the following software analysis protocol. Protein identification and quantification analysis were done with Integrated Proteomics Pipeline (www.integratedproteomics.com/). Tandem mass spectra were extracted into ms2 files from raw files using RawExtract 1.9.9<sup>32</sup> and were searched using ProLuCID algorithm<sup>33</sup> against the *C. neoformans* H99 database from the Broad Institute

(http://www.broadinstitute.org/annotation/genome/cryptococcus\_neoformans\_b/MultiDownloads.html, downloaded on August 8, 2012). The peptide mass search tolerance was set to 3 Da, and carboxymethylation (+57.02146 Da) of cysteine was considered to be a static modification. ProLuCID results were assembled and filtered using the DTASelect program<sup>34</sup> using two SEQUEST<sup>35</sup>-defined parameters, the cross-correlation score (XCorr) and normalized difference in cross-correlation scores (DeltaCN), to achieve a false-positive rate (1%). The following parameters were used to filter the peptide candidates: -p 1 -y 1 --trypstat --fpf 0.01 -in, where (-p) means minimum number of peptide per protein, (-y) minimum number of tryptic end per peptide, (--trypstat) statistics with tryptic status, (--fpf) false-positive rate for protein, and (-in) subset proteins included.

# **Data Analysis**

The software PatternLab<sup>36,37</sup> was used to identify differentially expressed and exclusive proteins found in biofilm and planktonic conditions. Pairwise comparisons between both conditions were performed using spectral count data on an updated version of the PatternLab's TFold module.<sup>38</sup> This module was used to select differentially expressed proteins under biofilm and planktonic conditions. The following parameters were used: proteins that were not detected in at least four out of six runs per condition were not considered, using a q-value of 0.05 and an F-stringency of 0.1. Lowabundance proteins, those with low p-values and a significant fold change but low spectral count values, were removed using the L-stringency of 0.4. Also, an absolute fold change greater than two was used to select differently expressed proteins. PatternLab's approximately area proportional Venn diagram (AAPV) module was used for pinpointing proteins uniquely identified under a condition using a probability 0.01.

The Blast2GO tool (http://www.blast2go.org)<sup>39</sup> was used to categorize the proteins detected by Gene Ontology (GO) annotation<sup>40</sup> according to biological process and molecular function. Additionally, the KEGG map module<sup>41</sup> allowed the display of enzymatic functions in the context of the metabolic pathways in which those proteins participate.

To investigate the characteristics of hypothetical proteins identified for biofilm, we used bioinformatics tools. The TargetP 1.0 (cutoff >0.9)<sup>42</sup> and TMHMM 2.0<sup>43</sup> were used to evaluate the subcellular location; SignalP 4.1<sup>44</sup> was used for prediction of secreted proteins (using cutoff default); ProtFun 2.2<sup>45</sup> was used for prediction of protein function, including cellular role, enzyme class, and Gene Ontology category; and HMMER 3.1<sup>46</sup> was used for searching sequence databases for homologues of protein sequences. TargetP, TMHMM, SignalP, and ProtFun programs are available at http://www.cbs.dtu.dk/services/; HMMER is available at http://hmmer.janelia.org/.

# **Interaction Data Set**

To construct an interface of our protein set with the known interactions for *Cryptococcus neoformans*, we downloaded from UniProt (www.uniprot.org) the file uniprot\_sprot\_fungi 01\_13 release. We extracted the FASTA sequences using an in-house software tool. We downloaded from STRING<sup>47</sup> the files proteins.links.detailed.v9.0.txt and protein.sequences.v9.0.fa.txt. We obtained the reference protein from STRING finding the closest sequence match to our protein set sequences.

Then, for every possible pair, an "ontological distance" 48 was calculated considering the fraction of GO terms shared over the

sum of distinct terms; the resulting score was then bounded between 0 and 1.

#### **Network Maps and Visualization**

We then parsed, using an in-house software tool, the STRING links file to extract the matching interaction. We then coupled the protein list with Gene Ontology<sup>40</sup> to gather the ontological terms. We calculated the frequency of each term in our list, and we clustered the proteins in groups based on their association with a particular term.

We developed an in-house software tool to integrate all of the different pieces of information to obtain a network interactions list. Each interaction is not directional and is represented by the corresponding score in the interaction data set. Each node in the interactions list was assigned a color based on its value in the protein regulation data set ranging in a gradient from blue, for down-regulated values, to red, for upregulated values. To visualize the network, we used the open-source Medusa viewer. <sup>49</sup>

### Validation of Proteomic Analysis

Catalase activity was assayed using hydrogen peroxide as substrate. <sup>50</sup> Phosphate buffer was added along with  $H_2O_2$  10 mM to 25 mL sample aliquots. Catalase activity was estimated by the decrease in absorbance of  $H_2O_2$  at 240 nm for 3 min. The decomposition of  $H_2O_2$  was followed at 240 nm (E=39.4 mm cm<sup>-1</sup>).

The assay for superoxide dismutase (SOD) was conducted as previously described. So A solution containing 0.05 M potassium phosphate buffer pH 7.8, 13 mM  $_{\rm L}$ -methionine, 75 mM NBT (nitro blue tetrazolium), 0.1 mM EDTA and 0.025% Triton X-100 was added to glass tubes. To start the reactions, we added the sample and 10 mM riboflavin at the same time that tubes were placed under fluorescent light for 15 min. After this period, absorbance was determined at 560 nm. SOD unit was defined by NBT reduction per mL  $h^{-1}$ .

Alanine aminotransferase assay was performed using TGP transaminase kit according to manufacturer instructions (Bioclin, Brazil).

Peptidase activity assays were tested upon chromogenic substrates: SF-17 (Bz-Phe-Val-Arg- $\rho$ NA; chymotrypsin/trypsin substrate), S2238 (H-D-Phe-Pip-Arg- $\rho$ NA; thrombin substrate), and S2251 (H-D-Val-Leu-Lys- $\rho$ NA; plasmin substrate), as previously described. Ten microliters of samples was incubated in 20 mM Tris-HCl buffer, pH 7.4. The reactions were initiated by adding substrates at 0.2 mM (final concentration) in a final volume of 100  $\mu$ L. Kinetic assays were monitored at 37 °C for 30 min in a SpectraMax spectrophotometer equipped with thermostat and shaking systems. One protease unit (U) was defined as the amount of enzyme that produces one  $\rho$ mol of  $\rho$ -nitroaniline per min per protein under the assay conditions described.

Phosphatase activity was measured by the rate of  $\rho$ -nitrophenol ( $\rho$ -NP) production. Samples (10  $\mu$ L) were incubated for 60 min at room temperature in 0.2 mL of reaction mixture containing 116.0 mM NaCl, 5.4 mM KCl, 30.0 mM HEPES-Tris buffer pH 7.0, and 5.0 mM  $\rho$ -nitrophenylphosphate ( $\rho$ -NPP) as substrate. The reaction was stopped by the addition of 0.2 mL of 20% trichloroacetic acid. Subsequently, the reaction mixture was centrifuged at 1500g for 15 min at 25 °C. The absorbance was measured spectrophotometrically at 405 nm using a microplate reader SpectraMax (Molecular Devices, USA). The concentration of released  $\rho$ -

nitrophenolate in the reaction was determined using a standard curve of  $\rho$ -nitrophenolate for comparison.

Western blot was also used to validate the proteomic data. In brief,  $20~\mu g$  of *C. neoformans* biofilm and planktonic proteins was separated by 4–20% Bis-tris gel (Invitrogen, CA) and transferred to a nitrocellulose membrane (Millipore, MA). Heat shock protein 70 (HSP70) (70 kDa) was detected with monoclonal antimouse antibody (1:1000 dilution). The secondary antibody (1:10.000) conjugated with HRP was detected using the SuperSignal West Pico chemiluminescent substrate (Thermo, IL).

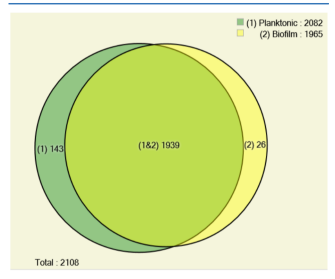
# **Statistical Analysis**

All enzymatic assays were performed in triplicate. Data generated from enzymatic activities were analyzed statistically using the Student's *t* test and GraphPad Prism 5 software.

# RESULTS

### **MudPIT Identification**

A total of 1965 proteins were identified from the biofilm and 2082 were identified from the planktonic condition according to maximum parsimony (Figure 1). The majority of those



**Figure 1.** Distribution of *Cryptococcus neoformans* H99 proteins obtained under biofilm and planktonic condition. Venn diagram shows the dispersion of total proteins identified under both conditions, according PatternLab's AAPV module, using 0.01 probability.

proteins were common to both conditions (1939), while 143 (6.78%) were identified only in planktonic cells and 26 (1.23%) were exclusive to cells growing in biofilm. All proteins identified in this research are listed in Table S1 and S2 (Supporting Information).

A total of 50 proteins were identified up- and 81 down-regulated in biofilm mode of growth (Table 1). The proteins that were up-regulated by more than two-fold included those related to oxidation—reduction (oxidoreductase), proteolysis (peptidases), and response to stress (catalase, glutathione-disulfide reductase, HSP70), and the down-regulated proteins were related to translation, transport and binding, transcription, and microtubule (Figure 2). We identified an aminotransferase with the highest differential expression (17.47 fold up-regulated) followed by lactoylglutathione lyase (15.4 fold), which is related to pyruvate metabolism and detoxification, and an uricase (15.28 fold) associated with uric acid degradation

(Table 1). The most down-regulated protein was aldehyde dehydrogenase (-51.09 fold), an enzyme responsible for the oxidation of aldehyde to carboxylic acid. Isocitrate lyase was down-regulated 26.75-fold and has function in the TCA cycle. Several ribosomal proteins were also down-regulated (Table 1).

Among the 26 proteins uniquely identified in biofilm, we identified proteins related to oxidative stress (Cu/ZN superoxide dismutase); acid phosphatase and two proteinases (Table 2). However, 46% of unique biofilm proteins were identified as conserved hypothetical, which were analyzed separately.

#### **Hypothetical Protein Analysis**

A total of 33 proteins identified were classified as hypothetical in the differentially expressed and unique protein sets in the biofilm. These proteins were studied further to identify possible function and subcellular localization. Their sequences were BLASTed against the NCBI nonredundant database, and all proteins had a high similarity with other hypothetical proteins. Therefore, we used other bioinformatic tools to identify subcellular localization or other possible molecular functions of the proteins (Table 3). According to ProtFun, we could identify proteins related to metabolism, biosynthesis, regulatory function, replication and transcription, purine/pyrimidine, and translation. In the down-regulated hypothetical proteins, 50% were identified as belonging to translation; for the up-regulated hypothetical proteins, regulatory function and metabolic process comprise 60%. This result agreed with previous reports, where biofilm cells are growing slower and thus have a lower translation rate. 21,22 According to HMMER program, proteins related to cytochrome c oxidase and RNA-recognizing motif were identified. Analyses of subcellular localization suggested that at least four proteins are present in or associated with the mitochondria (up-regulated and unique) and four proteins are possibly present in or associated with the transmembrane (only in biofilms) (Table 3). All proteins were considered unlikely to be secreted. The lack of similarity of these 33 pathogen-specific proteins with human proteins makes these good potential targets for therapeutic intervention.

# Kyoto Encyclopedia of Genes and Genomes (KEGG) Pathway Analysis

Recent works have indicated that bacterial biofilm cells have an active but altered metabolism. <sup>21,22,53</sup> To identify pathways affected by fungal biofilm formation in C. neoformans, we analyzed the MudPIT results using KEGG pathways in the Blast2Go program. Forty-three different pathways were associated with proteins identified as differentially regulated (up- or down-regulated) in the biofilm. The following KEGG pathways had four hits each: pyruvate metabolism, glycolysis/ gluconeogenesis, citrate cycle (TCA cycle), glyoxalate metabolism, and methane metabolism. Some pathways were represented only by up-regulated proteins: alanine, glutathione, nitrogen, sulfur, and tryptophan metabolisms. Analyzing the results in a wider view, we could observe that pathways related to the TCA cycle (succinate CoA ligase), glycolysis and pyruvate metabolism (aldehyde dehydrogenase, pyruvate carboxylase) are down-regulated. However, the second most up-regulated enzyme, lactoylglutathione lyase (pyruvate metabolism), is involved in detoxification and formation of glutathione and lactate, suggesting that pyruvate is probably being used in energy metabolism via lactate fermentation. (Figure S1, S2, and S3 in the Supporting Information). Analyzing other metabolic pathways related to amino acid metabolism (Figure S4 in the Supporting Information),

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accession number <sup>b</sup>	fold change <sup>c</sup>	p value	protein name
CNAG_02852T0	17.47	0.00094	aminotransferase
CNAG_04219T0	15.40	0.000119	lactoylglutathione lyase
CNAG_04307T0	15.28	0.012617	uricase
CNAG 01542T0	9.49	0.000858	taurine catabolism dioxygenase TauD
CNAG 03144T0	7.83	0.016609	protein-methionine-S-oxide reductase
CNAG 00457T0	6.71	0.000821	glutamine synthetase
CNAG_02714T0	6.63	0.008066	elongation factor 1-beta
CNAG_02399T0	6.55	$1.46 \times 10^{-05}$	glutathione-disulfide reductase
CNAG_00848T0	5.80	0.001169	conserved hypothetical protein
CNAG_02754T0	5.53	0.049673	40S ribosomal protein S12
	5	0.018703	conserved hypothetical protein
CNAG_03677T0	4.91	0.013703	conserved hypothetical protein
CNAG_07559T0			ThiJ/PfpI
CNAG_01686T0	4.54	0.000938	• •
CNAG_03482T0	4.38	0.013805	thioredoxin-dependent peroxide reductase
CNAG_04269T0	4.37	0.008231	peptidase
CNAG_05638T0	4.32	0.039603	conserved hypothetical protein
CNAG_05256T0	4.25	0.029929	catalase
CNAG_01375T0	4.08	0.004133	conserved hypothetical protein
CNAG_03509T0	4.04	0.022818	pyruvate dehydrogenase protein X component
CNAG_02202T0	3.84	0.021764	adenylyl-sulfate kinase
CNAG_04799T0	3.64	0.035322	ribosomal protein L14
CNAG_06302T0	3.53	0.012183	PEP2
CNAG_02076T0	3.49	0.002241	leukotriene-A4 hydrolase
CNAG_01577T0	3.48	0.005951	glutamate dehydrogenase
CNAG_07801T0	3.36	0.010318	aldo-keto reductase
CNAG_06443T0	3.22	0.042654	heat shock protein 70
CNAG_01558T0	3.14	0.004449	zinc-binding dehydrogenase
CNAG_07522T0	3.08	0.001507	conserved hypothetical protein
CNAG_04760T0	3	0.007741	cytoplasmic protein
CNAG_07771T0	3	0.003527	peptidase
CNAG_02568T0	2.94	0.005715	UBA/TS-N domain-containing protein
CNAG_01341T0	2.92	0.037823	mannose-6-phosphate isomerase
CNAG_06226T0	2.92	0.027042	conserved hypothetical protein
CNAG_02099T0	2.90	0.00182	fatty acid synthase beta subunit
CNAG_07362T0	2.79	0.015629	single-stranded DNA binding protein
CNAG_04441T0	2.66	0.006413	polyadenylate-binding protein
CNAG_01168T0	2.64	0.000321	centromere/microtubule binding protein cbf5
CNAG_00238T0	2.64	0.020937	isoleucine-tRNA ligase
CNAG_04604T0	2.53	0.032385	tryptophan-tRNA ligase
CNAG_01557T0	2.5	0.039665	calmodulin 1b
CNAG_03705T0	2.48	0.024666	conserved hypothetical protein
CNAG_01138T0	2.48	0.028101	cytochrome c peroxidase
CNAG_02726T0	2.39	0.041969	cytoplasmic protein
CNAG_00483T0	2.33	0.017333	actin
CNAG_02843T0	2.28	0.034824	conserved hypothetical protein
CNAG_05521T0	2.2	0.022853	aldose reductase
CNAG_05602T0	2.19	0.010051	1-pyrroline-5-carboxylate dehydrogenase
CNAG_03463T0	2.17	0.011508	conserved hypothetical protein
CNAG_00100T0	2.16	0.015629	chaperone
CNAG_01102T0	2.11	0.016887	oxidoreductase
CNAG_05900T0	-2.07	0.01048	glycine-tRNA ligase
CNAG_04209T0	-2.17	0.017508	voltage-gated potassium channel beta-2 subunit
CNAG_04601T0	-2.22	0.021853	glycine hydroxymethyltransferase
CNAG_03320T0	-2.26	0.014593	mannose-1-phosphate guanylyltransferase
CNAG_04904T0	-2.3	0.029246	clathrin heavy chain 1
<del>-</del>	-2.38	0.019719	heterogeneous nuclear ribonucleoprotein HRP1
 CNAG_06474T0			
	-2.39	0.007817	fatty-acid synthase complex protein
CNAG_06474T0		0.007817 0.005589	fatty-acid synthase complex protein translation initiation factor
CNAG_06474T0 CNAG_02100T0	-2.39		

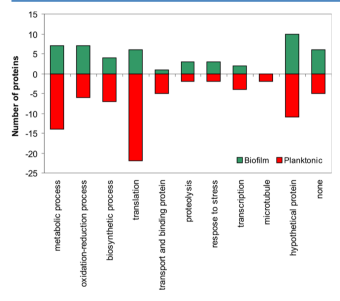
Table 1. continued

accession number <sup>b</sup>	fold change <sup>c</sup>	p value	protein name
CNAG_06900T0	-2.61	0.002331	phosphoglycerate mutase
CNAG_06866T0	-2.64	0.000455	conserved hypothetical protein
CNAG_01839T0	-2.71	0.046983	transcriptional elongation regulator
CNAG_04009T0	-2.72	0.005826	leucyl aminopeptidase
CNAG_06935T0	-2.73	0.025741	isochorismatase hydrolase
CNAG 01480T0	-2.77	0.00208	60S ribosomal protein L12
CNAG_03007T0	-2.78	0.000187	conserved hypothetical protein
CNAG_06113T0	-2.85	0.021278	conserved hypothetical protein
CNAG_05689T0	-2.90	0.029962	conserved hypothetical protein
CNAG_06511T0	-2.96	0.012872	conserved hypothetical protein
<del>-</del>	-3.12	0.037097	5-methyltetrahydropteroyltriglutamate-homocysteine S-methyltransferase
CNAG_01890T0	-3.12 -3.17		leucine-tRNA ligase
CNAG_06123T0		0.011257	e e e e e e e e e e e e e e e e e e e
CNAG_02489T0	-3.19	0.012336	mannitol-1-phosphate dehydrogenase
CNAG_02903T0	-3.21	0.037052	zinc-type alcohol dehydrogenase
CNAG_04105T0	-3.21	0.00296	hypothetical protein
CNAG_06231T0	-3.3	0.001301	ribosomal protein L13A
CNAG_00370T0	-3.33	0.011699	ubiquitin-carboxy extension protein fusion
CNAG_05907T0	-3.37	$2.19 \times 10^{-05}$	pyruvate carboxylase
CNAG_00565T0	-3.4	0.000322	VpsA
CNAG_00130T0	-3.61	0.000951	CAMK/CAMK1 protein kinase
CNAG_06150T0	-3.61	0.000726	heat-shock protein 90
CNAG_05886T0	-3.63	0.006982	ubiquitin conjugating enzyme MmsB
CNAG_04621T0	-3.68	0.001533	glycogen synthase
CNAG_01164T0	-3.75	0.030316	glutamine-fructose-6-phosphate transaminase
CNAG_03787T0	-3.78	0.021767	alpha tubulin
CNAG_00797T0	-3.79	0.021688	acetate-CoA ligase
CNAG_05555T0	-3.85	0.00402	ribosomal protein L4
CNAG_02335T0	-3.91	0.002194	UPF0364 protein
CNAG_00393T0	-4	0.013943	1,4-alpha-glucan-branching enzyme
CNAG_07346T0	-4	0.003779	t-complex protein 1
CNAG_05650T0	-4.02	0.015088	ubiquitin carboxyl-terminal hydrolase 5
CNAG_04652T0	-4.02	0.018038	enoyl reductase
CNAG 00147T0	-4.06	0.03385	splicing factor Prp8
CNAG 02118T0	-4.12	0.005713	conserved hypothetical protein
CNAG_04388T0	-4.66	0.000207	mitochondrial superoxide dismutase Sod2
CNAG_03641T0	-4.67	0.008395	eukaryotic translation initiation factor 3 subunit 7
CNAG 06235T0	-4.70	0.002397	conserved hypothetical protein
CNAG_04640T0	-4.88	$9.29 \times 10^{-05}$	acyl protein
CNAG_06125T0	-4.88	$4.94 \times 10^{-05}$	translation elongation factor 1 alpha
CNAG 00821T0	-5.06	0.031574	60s ribosomal protein 134-b
_	-5.21		predna95
CNAG_03780T0		0.001024	-
CNAG_00640T0	-5.49 -5.63	0.024398	40s ribosomal protein endoribonuclease L-PSP
CNAG_04066T0	-5.63	0.040426	
CNAG_01884T0	-5.86	0.012792	large subunit ribosomal protein L3
CNAG_02585T0	-6.7	0.013335	RfeF
CNAG_07316T0	-6.82	0.00049	alcohol dehydrogenase
CNAG_01870T0	-6.82	0.005086	mitochondrial protein
CNAG_07347T0	-6.93	0.000969	heat shock protein
CNAG_01486T0	-7.48 	0.011999	60S ribosomal protein L1Sb
CNAG_07746T0	-7.52	0.007699	methylenetetrahydrofolate dehydrogenase (NADP)
CNAG_04068T0	-7.53	0.014164	60s ribosomal protein
CNAG_00678T0	-7.67	0.002351	urease accessory protein ureG
CNAG_03010T0	-7.76	0.000503	enoyl-CoA hydratase/isomerase family protein
CNAG_01794T0	-7.93	0.015624	2-hydroxyacid dehydrogenase
CNAG_04445T0	-8.34	0.013355	40S ribosomal protein S7
CNAG_02129T0	-9.13	0.000584	conserved hypothetical protein
CNAG_05251T0	-9.33	0.000166	conserved hypothetical protein
CNAG_03048T0	-9.73	0.047775	cytoplasmic protein
 CNAG_07862T0	-10	0.000122	fumarate reductase
CNAG 01840T0	-11.10	0.03059	beta1-tubulin

Table 1. continued

accession number <sup>b</sup>	fold change <sup>c</sup>	p value	protein name
CNAG_00012T0	-13.08	0.012064	conserved hypothetical protein
CNAG_00747T0	-13.64	$3.23 \times 10^{-05}$	succinate-CoA ligase
CNAG_01224T0	-14.48	0.002602	ribosomal protein L18.e
CNAG_06605T0	-16.25	0.006007	ribosomal protein S2
CNAG_05721T0	-16.95	0.004036	peroxisomal hydratase-dehydrogenase-epimerase
CNAG_00116T0	-18.89	$1.00 \times 10^{-05}$	ribosomal protein S3
CNAG_00490T0	-21.43	0.006986	acetyl-CoA C-acyltransferase
CNAG_05303T0	-26.75	0.000987	isocitrate lyase
CNAG_00656T0	-32.32	$1.00 \times 10^{-05}$	60s ribosomal protein 17
CNAG_06628T0	-51.09	0.00014	aldehyde dehydrogenase

"Proteins listed in this Table were found to be statistically differentially expressed using PatternLab's Tfold module with an absolute fold change greater than 2.0 (BH-FDR 0.05). "According to Broad Institute ID. "Based on spectral count numbers obtained from biofilm and planktonic. Negative numbers represent down-regulated proteins in biofilm compared with planktonic condition.



**Figure 2.** Plot of the biological processes classification of all up- and down-regulated proteins in the biofilm of *Cryptococcus neoformans* H99 compared with the planktonic mode of growth. All of the tabulated proteins met a two-fold cutoff threshold for differential expression.

glutamine could be accumulated in biofilm. This compound could be used as a precursor for energy production. Figure 3 shows a schematic model proposed by *C. neoformans* biofilm for energy acquisition. The pathway related to glutathione metabolism (Figure S5 in the Supporting Information) shows that several enzymes are up-regulated, suggesting glutathione accumulation, which could be used for protection. The similarity of bacterial and fungal metabolic changes in biofilm growth hints at a common survival mechanism used by these distinct microbes.

#### **Validation**

To validate the proteomic analysis, we performed enzymatic assays and Western blot. Alanine aminotransferase activity (Figure 4A) was measured to be increased 15-fold, consistent with the 17-fold difference in spectral counts. Similarly, catalase activity was increased seven-fold, with a four-fold difference in spectral counts (Figure 4B). The activity of two proteins (superoxide dismutase and phosphatase) uniquely identified in biofilm was measured. As expected, more activity was observed in the biofilm (Figure 4C,D).

Protease assays were used to validate some measurements in the proteomic data. Multiple proteins identified as proteases were more abundant (at least three-fold) in biofilm (Table 1). To test cumulative protease activity, we measured the activity of lysates against three different synthetic substrates: related to coagulation (S2238), plasminogen degradation (S2251) and trypsin-like protease (SF-17). The activity was higher when *C. neoformans* was grown in biofilm as compared with planktonic, varying from 1.5- to 2.5-fold increase (Table 4) for all substrates evaluated consistent with the differences detected in protein level.

Western blot was used to validate the HSP70 protein. As observed in the Figure S6 in the Supporting Information, the immunoblot showed greater detection in *C. neoformans* biofilm as compared with planktonic, with an increase of almost two-fold according to ImageJ program (data not shown).

#### **Protein Interaction Networks**

To evaluate the network of those proteins identified uniquely and differentially expressed in the biofilm state, we generated protein interaction networks. Considering 157 proteins differentially expressed in biofilm, the interactome analysis revealed 105 protein-protein interactions (Figure 5). We observed that one protein represents a majority of connections: ribosomal protein L8 (CNAG\_05323T0). This protein was identified down-regulated in the biofilm state and could interact with 37 other proteins, most of them down-regulated (Figure 5B). Among these 37 possible connections, several are ribosomal proteins and some are related to metabolic processes. Evaluating the network by score, that is, closest similarity in the ontological distance, we identified one important group, where the majority of proteins were related to ribosome assembly (Figure 5C). Two other proteins with high interaction scores were Cu/Zn SOD (CNAG 01019T0) and SOD 2 mitochondrial (CNAG 04388T0).

# DISCUSSION

This is the first report evaluating the yeast *C. neoformans* biofilm using proteomics. Among the almost 2000 proteins detected, we found 76 proteins that were unique or more abundant in the biofilm state. Surprisingly, many (33) are uncharacterized hypothetical proteins. Many others are key metabolic enzymes that were previously identified in studies of bacterial biofilms. Together, these proteins provide potential and interesting targets for further studies.

Table 2. Unique Proteins Identified in C. neoformans Biofilm

accession number	protein name	spec count
CNAG_01019T0	Cu/Zn superoxide dismutase	109
CNAG_04236T0	acid phosphatase	42
CNAG_01147T0	ARP2/3 complex 20 kDa subunit	33
CNAG_04687T0	stearoyl-CoA 9-desaturase	27
CNAG_00351T0	predicted protein	23
CNAG_00668T0	conserved hypothetical protein	21
CNAG_02739T0	conserved hypothetical protein	19
CNAG_03926T0	methionine aminopeptidase 1	19
CNAG_06274T0	hypothetical protein	16
CNAG_04914T0	conserved hypothetical protein	13
CNAG_02914T0	conserved hypothetical protein	12
CNAG_03589T0	adrenodoxin-type ferredoxin	12
CNAG_04735T0	extracellular elastinolytic metalloproteinase	12
CNAG_06589T0	endoribonuclease L-PSP	11
CNAG_01650T0	60s ribosomal protein 17	10
CNAG_01991T0	cytochrome c oxidase subunit V	10
CNAG_04032T0	ATPase	10
CNAG_01381T0	conserved hypothetical protein	9
CNAG_03361T0	conserved hypothetical protein	9
CNAG_04900T0	WD40 protein Ciao1 variant	9
CNAG_06749T0	conserved hypothetical protein	9
CNAG_03228T0	universal stress protein family domain-containing protein	8
CNAG_05573T0	conserved hypothetical protein	8
CNAG_04257T0	conserved hypothetical protein	7
CNAG_06817T0	UAP1	7
CNAG_06716T0	conserved hypothetical protein	5

# Proteins Differentially Regulated in C. neoformans Biofilm

In our proteomic data, we were able to identify several proteins related to oxidative stress as differentially regulated or unique in the C. neoformans biofilm state: catalase, thioredoxin-dependent peroxide reductase, glutathione-disulfide reductase, Cu/Zn superoxide dismutase, oxidoreductase, and HSP70, among others. Activation of the oxidative stress response is one hypothesis that has been proposed to explain the increased resistance in microbial biofilms. It has gained considerable attention in recent years, with increasing evidence from several proteomic and transcriptomic published papers on bacterial and fungal biofilms. For example, HSP70, a protective protein related to thermal and oxidative stress, was also found to be upregulated in surface-associated proteins in the biofilm of another important pathogenic fungus, Candida albicans.<sup>25</sup> In another study, a greater number of stress-response proteins including SOD, thioredoxin, and HSP70 were up-regulated in Actinomyces naeslundii biofilm. 54 In microbial pathogens, HSPs are known to regulate virulence or immunological mechanisms.<sup>30</sup> For *C. neoformans*, this protein seems to act by additional mechanisms, influencing the interaction of yeasthost cells,<sup>55</sup> being considered the major antigen detected in patients with cryptococcosis. 56,57 In our interactome analysis, Cu/ZN SOD presented strongest interaction with other SOD, reinforcing the possible importance of these proteins in biofilms. In Shiga toxin-producing Escherichia coli, mutants lacking this enzyme were more susceptible to H2O2 than the wild type, displaying significant reduction in their capacity to adhere to epithelial cells and abiotic surfaces.<sup>58</sup> Catalase, an important enzyme in protecting the cell from oxidative damage by reactive oxygen species, was identified in biofilms and is thought to be related to antimicrobial resistance. 59-61 As expected, catalase activity was higher in biofilm cells than

planktonic, showing evidence of their participation in ROS protection in the biofilm. Interestingly, thioredoxin-dependent peroxidase reductase and catalase were identified in *C. albicans* and had the capacity to bind plasminogen. <sup>25,62</sup> We could identify a possible interaction between these proteins according to our interactome analysis.

Proteases could potentially help fungal dissemination into the host.<sup>63</sup> In our study, two proteases (methionine aminopeptidase and elastinolytic metalloproteinase) were identified uniquely in the biofilm. Extracellular proteolytic activity was linked to a role in the dissemination of cryptococcosis, being able to cleave key host components of the basement membrane and extracellular matrix, which may be a relevant factor in the fungal invasion. 65,65 Activation of plasminogen-to-plasmin is used by several microbial pathogens to increase their capacity for tissue invasion. 66,67 In C. neoformans, plasminogen can passively coat the surface of pathogens and, upon conversion to plasmin, facilitate pathogen dissemination by degrading vascular barriers.<sup>68</sup> Interestingly, protease activity on plasmin-specific substrate was higher in biofilm cells, which could link this resistant mode of growth to an infective behavior, where the cells would be prepared to invade the host tissue. Therefore, our hypothesis is that C. neoformans biofilm cells are more prepared to spread into the host.

Extracellular phosphatases have been implicated in the adhesion of several microbial pathogens and in host cell immune responses and play critical roles in signaling and biotic stress. S2,69,70 Similarly, we have seen an increase in phosphatase protein and activity in *C. neoformans* biofilm. However, our results are from cell lysate, and the phosphatase we identified may be acting either outside the cell altering the host environment or inside the cell in a regulatory fashion.

Table 3. Putative Classification and Localization of Cryptococcus neoformans H99 Hypothetical Proteins Identified As Differentially Regulated in Biofilms<sup>a</sup>

	HMMER			aldose 1-epimerase	ETC complex I subunit conserved region	dienelactone hydrolase family	RNA recognition motif	eisosome component PIL1	GDP/GTP exchange factor Sec2p	cofilin/tropomyosin-type actin-binding and SH3 domain	eisosome component PIL1				oxidoreductase family, NAD-binding	AdoMet dependent proline dimethyltransferase	short chain dehydrogenase			Stm1 and hyaluronan/mRNA binding family						cytochrome $c$ oxidase copper chaperone	mitochondrial large subunit ribosomal protein	cytochrome c oxidase copper chaperone			outer membrane protein TOM13	ARP2/3 complex 20 kDa subunit	fungal Zn(2)-Cys(6) binuclear cluster	3' exoribonuclease family, domain 1	LSM domain	RNA recognition motif	metallo-beta-lactamase superfamily
	ProtFun molecular function/enzyme class/GO		amino acid biosynthesis/-/-	amino acid biosynthesis/-/immune response	energy metabolism/-/growth factor	energy metabolism/lyase/-	metabolism/-/growth factor	regulatory function/-/growth factor	regulatory function/-/transcription regulation	regulatory function/-/transcription	translation/-/transcription	translation/ligase/growth factor		amino acid biosynthesis/lyase/growth factor	central intermediary metabolism/-/-	central intermediary metabolism/ligase/—	energy metabolism/lyase/-	replication and transcription/-/transcription regulation	translation/-/-	translation/-/growth factor	translation/-/transcription	translation/-/transcription regulation	translation/-/transcription regulation	translation/ligase/—		energy metabolism/-/transporter	energy metabolism/lyase/structural	fatty acid metabolism/—/transcription regulation	metabolism/lyase/immune response	purine and pyrimidine/—/growth factor	purine and pyrimidine/-/growth factor	regulatory function/-/growth factor	regulatory function/-/transcription regulation	translation/-/growth factor	translation/—/structural	translation/-/transcription	translation/ligase/-
,	TMHMM																									TM				$_{ m TM}$	TM			TM			
	signal P		Z	Z	Z	Z	Z	Z	Z	Z	Z	Z		Z	Z	Z	z	z	Z	z	Z	Z	Z	z		Z	Z	Z	Z	Z	Z	Z	Z	Z	Z	Z	Z
	target P				M						M																M								M		
	acc. number	Up-Regulated	$CNAG_03677T0$	CNAG_07559T0	CNAG_06226T0	CNAG_05638T0	CNAG_02843T0	CNAG_03705T0	CNAG_07522T0	CNAG_01375T0	CNAG_03463T0	CNAG_00848T0	Down-Regulated	CNAG_05689T0	CNAG_00012T0	CNAG_06235T0	CNAG_06511T0	CNAG_02118T0	CNAG_04105T0	CNAG_06113T0	CNAG_06866T0	CNAG_03007T0	CNAG_05251T0	CNAG_02129T0	Unique	CNAG_06274T0	CNAG_02914T0	CNAG_05573T0	CNAG_06749T0	CNAG_00668T0	CNAG_04257T0	CNAG_00351T0	CNAG_02739T0	CNAG_01381T0	CNAG_04914T0	CNAG_06716T0	CNAG_03361T0

<sup>a</sup>Target P: (M) mitochondrial; Signal P: (N) no peptide signal; TMHMM: (TM) transmembrane.

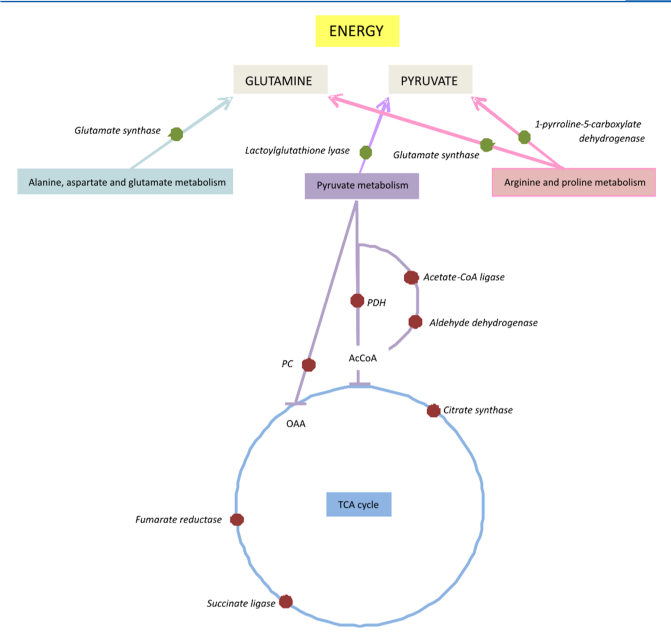


Figure 3. Schematic model of changes in metabolic regulation in biofilm cells of *Cryptococcus neoformans* H99 reveals a shift from the TCA cycle in energy acquisition. Red dots represent proteins down-regulated; green dots represent proteins up-regulated.

Another protein with a possible function in biofilm adhesion is alanine aminotransferase. This enzyme catalyzes the reversible reaction between pyruvate and glutamate and alanine and  $\alpha$ -ketoglutarate. It could be acting in pyruvate formation, which would be used for energy acquisition and therefore maintain the metabolic activity in biofilm cells. Otherwise, this protein is able to release alanine, an essential amino acid for cell maintenance. Alanine is present on *Enterococcus faecalis* and *Streptococcus* cell surfaces, and for these organisms, it is related to adhesion on surfaces and resistance. <sup>71,72</sup> It is not clear if this role for bacterial alanine aminotransferase is conserved in *C. neoformans*.

Although protein synthesis continued, many proteins related to assembly of ribosome subunits were down-regulated, consistent with reduced protein yield in biofilm. Corroborating this result, we observed several ribosomal proteins interacting in our network analysis. *C. gattii*, another species causing

cryptococcosis, presents down-regulation of ribosomal proteins in response to fluconazole, <sup>73</sup> showing that the maintenance of ribosome synthesis was affected due to the slowing of active metabolism and growth, consistent with our results. This evidence suggests that protein biosynthesis is affected in *C. neoformans* biofilms, as proposed for bacterial cells growing in biofilms. <sup>74–76</sup>

# **Hypothetical Proteins Analysis**

The differentially abundant proteins with known functions have been implicated in several cellular processes, including cell adhesion, host invasion, and metabolic regulation. However, a large portion of differentially abundant proteins have no known or theorized function.

Unique proteins identified in *C. neoformans* biofilm contain almost 50% of functionally unknown hypothetical proteins, suggesting that the complicated metabolic and regulation response to biofilm is still far from fully elucidated. Although, so

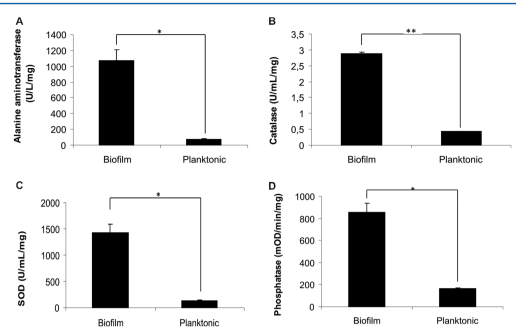


Figure 4. Enzymatic assays of (A) alanine aminotransferase, (B) catalase, (C) superoxide dismutase, and (D) phosphatase of biofilm and planktonic cells of *Cryptococcus neoformans* H99 (\*p < 0.001; \*\*p < 0.005).

Table 4. Peptidase Activities of Protein Extracts of Cryptococcus neoformans Cells Growing in Biofilm or Planktonic Condition against Three Different Chromogenic Protease Substrates<sup>a</sup>

substrate	biofilm/planktonic activity (pmol/min/mL/mg)
SF-17 (Bz-Phe-Val-Arg- $ ho$ NA)	2.5*
S2238 (H-D-Phe-Pip-Arg- $\rho$ NA)	2.5*
S2251 (H-D-Val-Leu-Lys- $\rho$ NA)	1.5**
a*p < 0.001, **p < 0.01.	

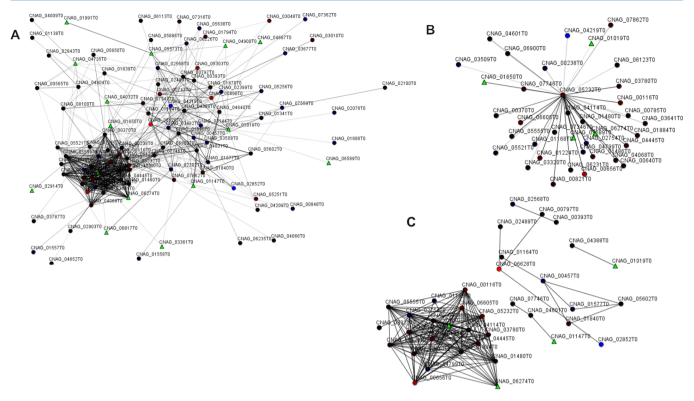
far, almost no functional information is available for these proteins. The goal here was to gain insights about localization and functioning for all hypothetical proteins identified. The bioinformatic analysis suggested down-regulated proteins related to translation and transcription, agreeing with our results compared with identified and categorized proteins. The role of the hypothetical proteins is not known but likely plays a role in the distinct physiological state of the biofilm, as described for *Desulfovibrio vulgaris*, a sulfate-reducing bacteria also implicated in a variety of human infections.<sup>77</sup> Although hypothetical proteins involved in biofilm changes have been reported before, further investigations are needed to evaluate their function because these proteins present homology to conserved hypothetical proteins from other microorganisms, including some pathogenic strains.

# Changes in Metabolic Profile of *C. neoformans* Biofilm – KEGG Analysis

A biofilm is a heterogeneous collection of cells with different metabolic states; the upper layers of a biofilm are more metabolically active cells, whereas in the lower layers the cells are in a state of quiescence. As well described for bacteria, the increased tolerance to antibiotics occurs when nutrients are limited. Therefore, slower rates of growth, changes in protein synthesis and metabolic activity present in lower layers can contribute to the increased persistence and consequent resistance of *C. neoformans* biofilms. The metabolic profiles for *C. neoformans* have shown that pyruvate level, related to energy metabolism, was higher in biofilm than in planktonic culture, indicating a reduction in energy production. Agreeing

with our results, a down-regulation in carbohydrate metabolism was observed in C. glabrata fungal biofilm, 80 and Staphylococcus mutans biofilms showed down-regulated glycolysis.<sup>81</sup> These changes in metabolic profiles may reflect reduced metabolic activity, which might contribute to persistence in biofilms. This is consistent with previous results that showed a slower growth rate for bacterial biofilms in comparison with planktonic cells.<sup>21,22</sup> Also, in biofilms, the deeper layers of cells is located undergrowth-limiting conditions, with anaerobic or microanaerobic environment. These cells could be supported by pyruvate fermentation, allowing their maintenance with no or little oxygen.<sup>82</sup> Alcohol dehydrogenase, an enzyme involved in pyruvate fermentation, was identified as down-regulated in C. neoformans biofilm. Similar result were found in C. albicans biofilm, where disruption of Adh1p significantly enhanced biofilm ex vivo and in vivo using engineered human oral mucosa and rat models.83 These results suggest that downregulation of alcohol dehydrogenase could be a critical determinant of biofilm formation. Therefore, our results strongly suggest that for C. neoformans, as described for Acinetobacter baumannii and Pseudomonas aeruginosa, the biofilm cells may depend on pyruvate fermentation for longterm survival.<sup>21,22</sup>

In addition to an oxygen-limiting environment, the biofilm cells are in constant stress — lack of nutrients, excess of waste products, and changes in pH gradient. In addition to the enzymes previously discussed that are involved in oxidative stress (catalase, HSP, SOD), our results show the lactolylhydrolase lyase or glyoxalase I is up-regulated in the biofilm. This



**Figure 5.** Interactome of differentially expressed proteins identified in *C. neoformans* H99 biofilms. Spheres and triangles represent proteins and lines connecting spheres indicate interactions between proteins. Red spheres are proteins down-regulated in biofilms; blue spheres indicate proteins upregulated; green triangles indicate unique proteins identified in biofilm. (A) General interactome; (B) cluster identifying proteins with higher connectivity; and (C) clusters showing higher score of strongest interactions.

enzyme is related to pyruvate metabolism, where it can also act in the detoxification of methylglyoxal, a side product of several metabolic pathways, releasing D-lactate and restoring glutathione. Glutathione is an important cellular antioxidant and could be used for protection in biofilm cells. However, further studies are needed to explore the real function of glyoxalase I in biofilms.

# CONCLUSIONS

C. neoformans biofilm formation on medical devices is considered a major source of foreign-body-associated infection. Through a shotgun proteomic approach comparing cells growing under biofilm and planktonic conditions, we provide insights about the metabolic mechanisms involved in cell maintenance, persistence and resistance. Giving special attention to those proteins differentially regulated in biofilms, this paper suggests a model of energy acquisition for biofilm cells. Although further investigations are needed, the fermentation pathway can be suggested as a target for new drugs. Within its biofilm, C. neoformans cells experience a highly stressful environment, with low oxygen and nutrient content and high waste products, showing a low rate of growth. To maintain the cells, pyruvate and glutamine appear to be contributing to growth and maintenance of the biofilm. Proteins involved in oxidative stress and related to tissue invasion were up-regulated, contributing to biofilm resistance and dispersion. These data suggest a metabolic correlation between bacterial and fungal biofilms. The proteins identified here provide additional information about the C. neoformans biofilm lifestyle. In conclusion, our proteomic data suggest general changes in metabolic processes, turnover, and global

stress responses. Understanding the pathways associated with biofilm maintenance and resistance is key to evaluating the effects of antifungal drugs, which are urgently needed.

# ASSOCIATED CONTENT

#### Supporting Information

Kegg pathway of the TCA cycle showing proteins down-regulated. Kegg pathway of the glycolysis/gluconeogenesis showing proteins downregulated. Kegg pathway of the pyruvate metabolism showing proteins down-regulated or up-regulated. Kegg pathway of the alanine, aspartate, and glutamate metabolism showing proteins up-regulated. Kegg pathway of the glutathione metabolism showing proteins up-regulated. Immunoblot of Cryptococcus neoformans biofilm and planktonic cells using monoclonal anti-mouse HSP70 antisera. This material is available free of charge via the Internet at http://pubs.acs.org.

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#### Notes

The authors declare no competing financial interest.

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