

The homeostasis of iron, copper, and zinc in *Paracoccidioides brasiliensis*, *Cryptococcus neoformans* var. *grubii*, and *Cryptococcus gattii*: a comparative analysis

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Célia Maria de Almeida Soares, Laboratório de Biologia Molecular, Departamento de Bioquímica e Biologia Molecular, Instituto de Ciências Biológicas, Universidade Federal de Goiás, ICB II, Campus II, 74690-900 Goiânia, Goiás, Brazil. e-mail: celia@icb.ufg.br Iron, copper, and zinc are essential for all living organisms. Moreover, the homeostasis of these metals is vital to microorganisms during pathogenic interactions with a host. Most pathogens have developed specific mechanisms for the uptake of micronutrients from their hosts in order to counteract the low availability of essential ions in infected tissues. We report here an analysis of genes potentially involved in iron, copper, and zinc uptake and homeostasis in the fungal pathogens Paracoccidioides brasiliensis, Cryptococcus neoformans var. grubii, and Cryptococcus gattii. Although prior studies have identified certain aspects of metal regulation in Cryptococcus species, little is known regarding the regulation of these elements in *P. brasiliensis*. We also present amino acid sequences analyses of deduced proteins in order to examine possible conserved domains. The genomic data reveals, for the first time, genes associated to iron, copper, and zinc assimilation and homeostasis in P. brasiliensis. Furthermore, analyses of the three fungal species identified homologs to genes associated with high-affinity uptake systems, vacuolar and mitochondrial iron storage, copper uptake and reduction, and zinc assimilation. However, homologs to genes involved in siderophore production were only found in P. brasiliensis. Interestingly, in silico analysis of the genomes of P. brasiliensis Pb01, Pb03, and Pb18 revealed significant differences in the presence and/or number of genes involved in metal homeostasis, such as in genes related to iron reduction and oxidation. The broad analyses of the genomes of P. brasiliensis, C. neoformans var. grubii, and C. gattii for genes involved in metal homeostasis provide important groundwork for numerous interesting future areas of investigation that are required in order to validate and explore the function of the identified genes and gene pathways.

Keywords: micronutrient homeostasis, pathogenic fungi, infection

INTRODUCTION

A sufficient supply of iron, copper and zinc is essential for all living and proliferating organisms. In infectious diseases, iron, copper and zinc metabolism are important for both the host and the pathogen, and complex responses in each occur to maintain adequate resources of these elements to preserve homeostasis. Iron, in the form of heme and iron–sulfur clusters, is essential as a cofactor of various enzymes, oxygen carriers, and electron-transfer systems involved in vital cellular functions ranging from respiration to DNA replication (Schaible and Kaufmann, 2004). Copper is a redoxactive metal ion essential for most aerobic organisms, which also serves as a catalytic and structural cofactor for enzymes involved in energy generation, iron acquisition, oxygen transport, and cellular metabolism, among other processes (Kim et al., 2008). Zinc is also a crucial metal, since it is at the catalytic center of numerous enzymes and plays important roles in the functionality of a wide variety of

proteins (Van Ho et al., 2002). Mammalian hosts and microbes have developed sophisticated strategies to acquire these metals, even under conditions in which their availability is limited. One of the strategies developed by mammalian hosts to prevent microbial infections is to limit the availability of iron (Weinberg, 2009). Recently, it has been demonstrated that zinc deprivation is a host defense mechanism utilized by macrophages during *Histoplasma capsulatum* infection (Winters et al., 2010). In addition, the binding of copper to calgranulin C in human neutrophils could be a mechanism of antimicrobial action (Moroz et al., 2003). In order to counteract these and other host responses, microorganisms employ a range of uptake mechanisms for the targeted acquisition of iron, copper and zinc.

Ferric iron is generally insoluble at physiological pH in the presence of oxygen. Thus, the common mechanisms of iron-assimilation include the reduction of ferric (Fe^{3+}) to ferrous (Fe^{2+}),

and solubilization of Fe³⁺ by binding siderophores (Kornitzer, 2009). The reductive system in fungi is regulated by three different mechanisms. First, a low-affinity iron reductase that functions in iron-rich environments generates Fe²⁺, which is transported into the cell by a non-specific low-affinity iron permease. Second, a regulated high-affinity ferric reductase operates in low iron conditions, such as those present in a mammalian host. The produced Fe²⁺ is further oxidized to Fe³⁺ by a membrane multi-copper-oxidase before being transported across the cell membrane by a high-affinity iron permease. The third mechanism is a non-enzymatic reduction, such as that promoted by 3-hydroxyanthranilic acid (3HAA), which is known to maintain a reduced environment to facilitate the release and sustain the presence of Fe²⁺ at the fungal membrane until transport occurs (Howard, 1999).

Ferric iron uptake mediated by siderophores is considered a non-reductive high-affinity mechanism by which microorganisms acquire iron. Siderophores are low-molecular weight ($M_r < 1500$), ferric iron-specific chelators (Neilands, 1993). Microorganisms produce siderophores as scavenging agents in low iron concentration environments in order to supply iron to the cell through the solubilization of extracellular ferric iron. Siderophores are also produced intracellularly for iron storage in most fungi (Matzanke et al., 1987). Siderophores can be classified into three main groups depending on the chemical nature of the moieties donating the oxygen ligands for Fe³⁺: catechols, carboxylates and hydroxamates (Miethke and Marahiel, 2007). With the exception of the carboxylate rhizoferrin produced by zygomycetes, the other known fungal siderophores are all hydroxamates (Van der Helm and Winkelmann, 1994). Fungal hydroxamates are derived from the non-proteinogenic amino acid ornithine and can be grouped into four structural families: rhodotorulic acid, ferrichromes, coprogens and fusarinines. Siderophores are named based on their iron-charged forms, existing in the iron-free form of the ligand called desferri-siderophore. Not all fungi produce siderophores. For example, Saccharomyces cerevisiae is not a siderophore producer (Neilands et al., 1987). Similarly, Cryptococcus species and Candida albicans are also unable to produce siderophores. However, these pathogenic fungi can utilize iron bound to siderophores secreted by other species (bacteria and fungi), the xenosiderophores (Howard, 1999). After siderophores are synthesized, they can be utilized intracellularly or secreted to the extracellular medium to solubilize ferric iron. For secreted siderophores, the captured metal of the siderophore-iron complex may be utilized either by reductive iron assimilatory systems or by internalization of the whole complex by specific transporters. In fungi, the uptake of siderophore-iron chelates is accomplished by transporters of the siderophore-iron transporter (SIT) subfamily, previously designated as family 16 of the major facilitator superfamily (MFS; Pao et al., 1998). These transporters are integral membrane proteins, with 12-14 predicted transmembrane domains, that mediate the import of siderophores in a highly regulated process (Philpott, 2006).

Several homeostatic mechanisms that ensure the maintenance of copper at a sufficient concentration for cell growth have been identified. Copper homeostasis in fungi is maintained by the transcriptional regulation of genes involved in copper acquisition, mobilization and sequestration and also at the posttranslational level (Gross et al., 2000). In *S. cerevisiae* copper is reduced from Cu (II) to Cu (I) by cell surface metalloreductases (Hassett and Kosman, 1995; Georgatsou et al., 1997) and uptake is mediated by Ctr1p and Ctr3p, two high-affinity transporters. Both *ctr1* and *ctr3* genes are regulated at the transcriptional level in response to copper availability, being induced by copper deprivation (Dancis et al., 1994a; Pena et al., 2000). The vacuolar copper transporter Ctr2p is also involved in the intracellular copper homeostasis, since it provides copper via mobilization of intracellular copper stores (Rees et al., 2004).

Zinc homeostasis is maintained by posttranslational and transcriptional homeostatic regulatory mechanisms (Lyons et al., 2000; Eide, 2003). Unlike iron and copper, zinc is taken up as divalent cation. Once inside the cell, zinc is neither oxidized nor reduced (Berg and Shi, 1996). In S. cerevisiae the uptake of zinc is mediated by two separate systems. One system has a high-affinity for this metal and is active in zinc-limited conditions (Zhao and Eide, 1996a). The second system has a lower affinity for zinc and is not highly regulated by zinc concentrations (Zhao and Eide, 1996b). The expression of the high-affinity zinc transporter Zrt1p and the low-affinity zinc transporter Zrt2p is regulated by the transcription factor Zap1p, which plays a central role in zinc homeostasis (Zhao and Eide, 1997). The zinc transporter activity is also posttranslationally regulated. High levels of extracellular zinc trigger the inactivation of Zrt1p through endocytosis of the protein and its subsequent degradation in the vacuole (Gitan et al., 1998).

This paper focuses on the metabolism of iron, copper and zinc in the fungal pathogens Paracoccidioides brasiliensis, Cryptococcus neoformans var. grubii, and Cryptococcus gattii. Low iron conditions have been associated with the susceptibility of P. brasiliensis, the etiological agent of paracoccidioidomycosis (PCM), to the antimicrobial action of monocytes (Dias-Melicio et al., 2005). Major phenotypic changes in C. neoformans, the etiological agent of cryptococcosis, are regulated by iron availability. For example, low iron concentrations result in the induction of capsule enlargement and the repression of laccase (Jung and Kronstad, 2008). Although iron regulation is well described in Cryptococcus species (Jung et al., 2008), iron associated processes are poorly understood in P. brasiliensis. Further, there is limited information on the impact of copper and zinc in *P. brasiliensis*, as well as the impact of zinc in Cryptococcus species. In this paper we performed in silico analyses of genes related to iron, copper and zinc metabolism in P. brasiliensis, C. neoformans var. grubii and C. gattii. We also compared the obtained information with data available from S. cerevisiae, which represents the most deeply studied model fungus, and other fungi.

MATERIALS AND METHODS

Sequences of genes related to iron, copper and zinc uptake, as well as to siderophore biosynthesis and uptake were used in the search of orthologs of *P. brasiliensis* and *Cryptococcus* species genomes. The *P. brasiliensis* database¹ includes the genomes of three isolates (*Pb*01, *Pb*03, and *Pb*18) and the cryptococcal database includes genomes of *C. neoformans* var. *grubii*² and *C. gattii*³. The sequences used in

¹http://www.broadinstitute.org/annotation/genome/paracoccidioides_brasiliensis/ MultiHome.html

²http://www.broadinstitute.org/annotation/genome/cryptococcus_neoformans/ MultiHome.html

³http://www.broadinstitute.org/annotation/genome/cryptococcus_neoformans_b/ MultiHome.html

the *in silico* analysis were obtained from the NCBI databank⁴, and they are primarily from *S. cerevisiae*, but also include genes from other fungi, such as *Aspergillus fumigatus*, *Aspergillus nidulans*, *C. albicans* and *H. capsulatum*. The search by orthologs was based on sequence similarity by using the BLAST tool. The expectation value adopted in the databases search was *E*-value $\leq 10^{-5}$.

The deduced amino acid sequences of the orthologs found in *P. brasiliensis* isolates and *Cryptococcus* species were analyzed. Searches for conserved domains and signal peptides in the orthologs proteins were performed using the Conserved Domain Database at NCBI⁵ and the online software SMART⁶. Predictions of putative transmembrane segments were made using the TopPred⁷ server and SMART software. Amino acid sequences alignment were performed using the ClustalX2 (Larkin et al., 2007).

RESULTS AND DISCUSSION IRON

Uptake of iron at the cell surface by the reductive system

To better understand how P. brasiliensis could acquire iron by the reductive system, in silico analyses were performed utilizing S. cerevisiae⁸ and C. albicans⁹ sequences. The data showed that Pb01 contains four metalloreductase (Frep) homologs, Pb03 five homologs, and Pb18 three homologs (Table 1). The genes encoding metalloreductases were fre1, fre3, fre5, fre7 and frp1. Also, Pb01 and Pb03 have two homologs each of the ferroxidase Fetp and Pb18 has one. The reductive uptake system was first described in S. cerevisiae (Lesuisse et al., 1987). The enzymatic reduction step in S. cerevisiae is catalyzed by members of the FRE family of metalloreductases. The products of the *fre* genes are not specific for iron reduction, since they can also promote copper reduction. S. cerevisiae Fre1p and Fre2p are required for growth on media with low concentrations of ferric iron salts. Fre3p and Fre4p catalyze uptake of iron from siderophores and Fre7p is under the control of the copperdependent transcription factor Mac1p (Philpott and Protchenko, 2008). The expression of C. albicans ferric reductase Frp1p is upregulated by alkaline pH and iron-limited conditions (Liang et al., 2009). Future studies are required to dissect the roles of the different P. brasiliensis reductases, especially in in vivo conditions.

Homologs for iron permeases (Ftrp and Fthp) were not found in *P. brasiliensis* genomes, corroborating the hypothesis that iron is transported by the zinc permeases, as previously suggested by transcriptional analyses (Bailão et al., 2006, 2007; Costa et al., 2007). However, in the present *in silico* analysis, we identified five zinc transporters (**Table 1**). These permeases could be coupled with one or more of the ferroxidases homologs (Fet5p, Fet31p and Fet33p) identified in the *P. brasiliensis* genome database. In *S. cerevisiae*, reduced iron is taken up through a high-affinity transport complex that consists of Fet3p, a multi-copper ferroxidase, and Ftr1p, a permease. Independent studies have demonstrated that Fet3p produced by *S. cerevisiae* $\Delta ftr1$ mutant cells is retained in a cytoplasmic compartment in a copper-free, inactive form. Correspondingly, Ftr1p produced by *S. cerevisiae* $\Delta fet3$ mutant cells fails to reach the plasma membrane (Stearman et al., 1996). These observations are in agreement with a model in which the two proteins form a heterodimer or higher order structure for correct maturation and trafficking to the plasma membrane (Kosman, 2003).

The *P. brasiliensis* genomes analysis revealed the presence of a *ggt1* homolog. This gene is presumably responsible for the glutathione (GSH)-dependent iron reduction activity previously identified in functional studies (Zarnowski and Woods, 2005). The proposed mechanism comprises secretion of a glutathione-dependent ferric reductase (GSH–FeR), named Ggt1p, that purportedly utilizes siderophores and Fe³⁺-binding proteins as substrates, enhancing the enzymatic activity under iron-limiting conditions, which is consistent with the function of a high-affinity uptake system, as described in *H. capsulatum* (Timmerman and Woods, 2001).

Homologs of permease genes involved in low-affinity iron reductive systems, such as *smf*, were not detected in our analysis. Hence, the low-affinity permease utilized by P. brasiliensis to acquire iron could be one of the zinc permeases, as suggested (Table 1). Despite the absence of iron permease fth1 gene homologs, P. brasiliensis has one ccc1 gene homolog that could drive iron vacuolar transport. P. brasiliensis also has homologs of the mitochondrial iron transporters genes mrs3 and mrs4 and the mitochondrial iron chaperone Yfh1p, suggesting mitochondrial iron homeostasis in this pathogen (Table 1). Since mitochondria are major users of iron, it follows that they should contain machinery required for its transport. Mrs3p and Mrs4p are homologous and functionally redundant proteins found in the inner mitochondrial membrane of S. cerevisiae, which are involved in transport under iron-limiting conditions (Foury and Roganti, 2002). Yfh1p, a homolog of human frataxin, is also involved in mitochondrial iron homeostasis (Babcock et al., 1997). While Mrs3p and Mrs4p mediate iron delivery from the outside to the inside of mitochondria, the frataxin homolog facilitates the use of iron within this organelle, functioning as a mitochondrial matrix iron chaperone (Zhang et al., 2006; Froschauer et al., 2009).

Cryptococcal genomic databases analysis revealed both S. cerevisiae and C. albicans homologs for proteins related to iron metabolism (Table 1). Remarkably, the C. neoformans var. grubii database contains four metalloreductase homologs, while the C. gattii genome has three similar homologs. The reason for the multiplicity of metalloreductases isoenzymes is not clear, although it is speculated that some sets of genes are expressed under specific conditions for iron acquisition (Kornitzer, 2009). Concerning the ferroxidases, C. neoformans var. grubii has three homologs and C. gatti contains one. Both genomes possess two iron permeases homologs, whose presence is supported by prior functional analyses (Jung et al., 2008). Two iron permeases, gene orthologs of S. cerevisiae ftr1, have been identified in C. neoformans, namely Cft1p and Cft2p (Jung et al., 2008). The expression of the *cft1* gene is down-regulated at high iron concentrations, suggesting that its product functions as a high-affinity iron permease. The role of cft2 is still unclear, although it supposedly encodes a low-affinity iron permease or a vacuolar permease that could transport stored iron to the cytoplasm, similar to what occurs in S. cerevisiae with the iron permease Fth1p. One of the iron permeases here identified is probably a Fth1p homolog, which

⁴http://www.ncbi.nlm.nih.gov/guide/

⁵http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi

⁶http://smart.embl-heidelberg.de/

⁷http://mobyle.pasteur.fr/cgi-bin/portal.py?form=toppred

⁸http://www.yeastgenome.org/

⁹http://www.candidagenome.org/

Gene	Organism/accession number	Predicted function	Orthologs in <i>Pb</i> 01, 03 and 18 (accession numbers)†	<i>E</i> -value*	Orthologs in <i>Cryptococcus</i> species (accession numbers) [†]	E-value*
fre1	<i>S. cerevisiae</i> NP_013315	Metalloreductase	PAAG_05370.1 PABG_06003.1	e-22 e-19	Not identified	_
fre3	<i>S. cerevisiae</i> NP_015026	Metalloreductase	PAAG_02079.1 PABG_02329.1 PADG_00813.1	e-35 e-35 e-35	Not identified	-
fre5	<i>S. cerevisiae</i> NP_015029	Metalloreductase	PABG_07812.1	e-26	Not identified	-
fre7	<i>S. cerevisiae</i> NP_014489	Metalloreductase	PAAG_06164.1 PABG_06497.1 PADG_07957.1	0.0 0.0 0.0	CNAG_00876.2 CNBG_6082.2	e-37 e-37
fre8	<i>S. cerevisiae</i> NP_013148	Metalloreductase	Not identified	-	CNAG_07334.2 CNBG_2116.2	e-10 e-07
fre10	<i>C. albicans</i> XP_711543	Metalloreductase	Not identified	-	CNAG_06821.2 CNBG_5888.2	e-34 e-27
cfl4	<i>C. albicans</i> XP_715639	Metalloreductase	Not identified	-	CNAG_06524.2	e-32
frp1	<i>C. albicans</i> XP_713315	Metalloreductase	PAAG_04493.1 PABG_04278.1 PADG_04652.1	e-26 e-26 e-26	Not identified	-
fet3	<i>S. cerevisiae</i> NP_013774	Ferroxidase	Not identified	-	CNAG_06241.2	0.0
fet5	<i>S. cerevisiae</i> NP_116612	Ferroxidase	PABG_05667.1 PADG_05994.1	e-40 e-37	CNAG_07865.2 CNBG_4942.2	0.0 0.0
fet31	<i>C. albicans</i> XP_711263	Ferroxidase	PAAG_06004.1	e-39	CNAG_02958.2	0.0
fet33	<i>C. albicans</i> XP_711265	Ferroxidase	PAAG_00163.1 PABG_05183.1	e-33 e-33	Not identified	-
ftr1/ftr2	<i>C. albicans</i> XP_715020/ XP_715031	Iron permease	Not identified	-	CNAG_06242.2 CNBG_3602.2	0.0 0.0
smf1	<i>S. cerevisiae</i> NP_014519	Low-affinity permease	Not identified	-	CNAG_05640.2 CNBG_6162.2	0.0 0.0
fth1	<i>C. albicans</i> XP_723298	Vacuolar transporter	Not identified	-	CNAG_02959.2 CNBG_4943.2	0.0 0.0
ccc1	S. cerevisiae	Vacuolar	PAAG_07762.1	e-31	CNAG_05154.2	e-23
	NP_013321	transporter	PABG_00362.1 PADG_02775.1	e-31 e-31	CNBG_4540.2	e-18
mrs3/ mrs4	<i>S. cerevisiae</i> NP_012402/ NP_012978	Mitochondrial iron transporter	PAAG_05053.1 PABG_04509.1 PADG_04903.1	0.0 0.0 0.0	CNAG_02522.2 CNBG_4218.2	0.0 0.0
yfh1	<i>S. cerevisiae</i> NP_010163	Mitochondrial matrix iron chaperone	PAAG_02608.1 PABG_03095.1 PADG_01626.1	e-15 e-09 e-16	CNAG_05011.2 CNBG_4670.2	e-18 e-18

Table 1 | Orthologs to genes related to iron, copper and zinc uptake by reductive systems in *P. brasiliensis* and *Cryptococcus* species.

Table 1 | Continued

Gene	Organism/accession number	Predicted function	Orthologs in <i>Pb</i> 01, 03 and 18 (accession numbers)†	<i>E</i> -value*	Orthologs in <i>Cryptococcus</i> species (accession numbers)†	<i>E</i> -value [®]
ggt1	H. capsulatum	Secreted	PAAG_06130.1	0.0	CNAG_02888.2	0.0
	EGC49121	glutathione-	PABG_06527.1	0.0	CNBG_3537.2	0.0
		dependent ferric reductase	PADG_07986.1	0.0		
mac1	S. cerevisiae	Copper	PAAG_08210.1	e-5	CNAG_07724.2	e-7
	NP_013734	metalloregulatory transcription factor	PABG_07429.1	e-5	CNBG_2252.2	e-7
ctr3	S. cerevisiae	High-affinity	PAAG_05251.1	e-22	CNAG_00979.2	e-14
	NP_013515	copper	PABG_07607.1	e-21	CNBG_0560.2	e-14
		transporter of the plasma membrane	PADG_05084.1	e-21		
ctr1	S. cerevisiae	High-affinity	Not identified	-	Not identified	-
	NP_015449	copper transporter of the plasma membrane				
ctr2	S. cerevisiae	Putative	PABG_01536.1	e-14	CNAG_01872.2	e-13
	NP_012045	low-affinity copper transporter of the vacuolar membrane	PADG_04146.1	e-14		
atx1	S. cerevisiae	Cytosolic copper	PAAG_00326.1	e-12	CNAG_02434.2	e-10
alXI	NP_14140	metallochaperone	PABG_06615.1	e-12 e-12	CNAG_02434.2 CNBG_4136.2	e-10
	111_14140	Thetailochaperone	PADG_02352.1	e-12 e-12	CNDO_4130.2	6-11
ccc2	S. cerevisiae	Cu ²⁺ transporting	PAAG_07053.1	0.0	CNAG_06415.2	0.0
0002	NP_010556	P-type ATPase	PABG_03057.1	0.0	CNBG_5045.2	0.0
		r type An use	PADG 01582.1	0.0	01000_0040.2	0.0
cup1	<i>S. cerevisiae</i> NP_011920	Metallothionein	Not identified	_	Not identified	-
cup2	<i>S. cerevisiae</i> NP_011922	Metallothionein	Not identified	-	Not identified	_
sod1	S. cerevisiae	Cytosolic	PAAG_04164.1	0.0	CNAG_01019.2	0.0
	NP_012638	superoxide	PABG_03954.1	0.0	CNBG_0599.2	0.0
		dismutase	PADG_07418.1	0.0		
sod2	S. cerevisiae	Mitochondrial	PAAG_02725.1	0.0	CNAG_04388.2	0.0
	NP_011872	superoxide	PABG_03204.1	0.0	CNBG_2661.2	0.0
		dismutase	PADG_01755.1	0.0		
zrt1	S. cerevisiae	High-affinity zinc	PAAG_08727.1	0.0	CNAG_03398.2	e-40
	NP_011259	transporter of the plasma membrane	PABG_07725.1 PADG_08567.1	0.0	CNBG_2209.2	e-41
zrt2	S. cerevisiae	Low-affinity zinc	PAAG_03419.1	e-27	CNAG_00895.2	0.0
	NP_013231	transporter of	PABG_05498.1	e-26		
		the plasma membrane	PADG_06417.1	e-28		

Table 1 | Continued

Gene	Organism/accession number	Predicted function	Orthologs in <i>Pb</i> 01, 03 and 18 (accession numbers)†	<i>E</i> -value*	Orthologs in <i>Cryptococcus</i> species (accession numbers) [†]	<i>E</i> -value*
zrc1	<i>S. cerevisiae</i> NP_013970	Vacuolar membrane zinc transporter	PAAG_00702.1	e-41	Not identified	-
cot1	<i>S. cerevisiae</i> NP_014961	Vacuolar membrane zinc transporter	PAAG_07885.1 PABG_07467.1 PADG_08196.1	e-44 0.0 0.0	CNAG_02806.2 CNBG_3460.2	e-40 e-37
zrt3	<i>S. cerevisiae</i> NP_012746	Vacuolar membrane zinc transporter	PAAG_09074.1 PABG_04697.1 PADG_05322.1	e-23 e-22 e-23	Not identified	-
msc2	<i>S. cerevisiae</i> NP_010491	Cation diffusion facilitator protein of the endoplasmic reticulum and nucleus	PABG_07115.1 PADG_06381.1	e-40 e-40	CNAG_05394.2 CNBG_4458.2	e-23 e-24
zap1	<i>S. cerevisiae</i> NP_012479	Zinc-regulated transcription factor	PAAG_03645.1 PABG_03305.1 PADG_01870.1	e-20 e-18 e-24	CNAG_05392.2 CNBG_4460.2	e-40 e-28

*Similarities with E-values < 10⁻⁵ were considered significant.

Accession numbers: PAAG refers to Pb01; PABG refers to Pb03; PADG refers to Pb18; CNAG refers to C. neoformans var. grubii and CNBG refers to C. gattii.

is likely involved in vacuolar iron uptake. Moreover, we could identify iron transporter ccc1 gene homologs in the genome, suggesting that a vacuolar iron homeostasis system exists in Cryptococcus. Data mining revealed one homolog of the low-affinity gene smf family, confirming the presence of both high and low-affinity iron reductase systems, as described (Jacobson et al., 1998). The presence of mitochondrial mrs3, mrs4 and yfh1 gene homologs in C. neoformans var. grubii supports a mechanism for iron homeostasis (Nyhus and Jacobson, 1999; Jacobson et al., 2005). Additionally, our in silico analyses demonstrated that cryptococcal reductive systems are closely related to that of S. cerevisiae (Table 1). Although no activity for the enzyme glutathione-dependent ferric reductase had been reported in Cryptococcus, both genomes contain ggt1 homologs suggesting the presence of a GSH-FeR system. A comparative analysis of iron uptake by reductive systems in P. brasiliensis, C. neoformans var. grubii and C. gattii is depicted in Figure 1.

Conserved domains in proteins related to the reductive iron metabolism

Amino acid sequence analyses of orthologs proteins found in the *P. brasiliensis* isolates and *Cryptococcus* species may support the assumption of conserved functions. Searching for conserved domains in all the analyzed sequences (**Table A1** in Appendix) revealed that most of the *P. brasiliensis* and *Cryptococcus* deduced proteins codified by the genes related to reductive iron metabolism contain conserved domains related to specific functions. Regarding to metalloreductases, the presence of a ferric reductase domain and a FAD- and/or a NAD-binding domain can be essential for functional enzymatic activity, since they are responsible for electron donation, as described in other organisms (De Luca and Wood, 2000). A schematic diagram presenting the cited motifs in a metalloreductase Frep is shown in **Figure 2**. An HPFTXXS motif is believed to be a site for FAD-binding and a glycine-rich motif and a cysteine–glycine couple are thought to be involved in NADPH binding (Shatwell et al., 1996). As well, copper-oxidase domains are required for ferroxidase activity. *S. cerevisiae* Fet3p is a multi-copper-oxidase and, like other copper proteins, possesses three distinct types of Cu²⁺-binding sites. Oxidation of Fe²⁺ occurs at the type 1 copper site followed by the reduction of molecular oxygen to 2H₂O at the other two copper sites (Hassett et al., 1998; Kosman, 2003). The ferroxidases in the *P. brasiliensis* isolates and *Cryptococcus* species present such domain, suggesting they are functional proteins.

Siderophore production

Culture supernatants of P. brasiliensis grown in media with low iron concentrations display higher iron binding capacity when compared with culture supernatants from iron-rich media (Arango and Restrepo, 1988), which has suggested that siderophores are involved in iron acquisition in this fungus. Furthermore, in silico analysis of P. brasiliensis structural genomes indicates that this fungus can potentially produce siderophores. The three sequenced P. brasiliensis genomes show sequences that potentially encode all the necessary enzymes for siderophore synthesis: sidA, sidF, sidC and sidD (A. fumigatus orthologs), as shown in Table 2 and Figure 1. This biosynthetic pathway may lead to the production of hydroxamate-type siderophores. The first committed step in siderophore biosynthesis is the N5-hydroxylation of ornithine catalyzed by ornithine-N5-oxygenase. The sid1 gene of Ustilago maydis, the etiologic agent of corn smut, was the first characterized fungal ornithine-N⁵-oxygenase-encoding gene (Mei et al., 1993).



Orthologs of *sid1* have been identified in *A. fumigatus (sidA)* and *H. capsulatum (sid1)*. In the latter, disruption of *sid1* causes poor growth under low iron conditions and loss of siderophore production, suggesting an important role of siderophore production in

iron-limiting conditions (Schrettl et al., 2004; Hwang et al., 2008). The formation of the hydroxamate group consists of the transfer of an acyl group from acyl-coenzyme A to N^5 -hydroxyornithine. Different acyl group usage results in the production of distinct



FIGURE 2 | Domains found in a Frep related to iron reductive uptake in *P*: *brasiliensis* isolates and *Cryptococcus* species. The found domains are: ferric reductase domain (black boxes), FAD-binding domain (dark gray boxes) and NAD-binding domain (light gray boxes). The length of each protein, in amino acids, is shown on the right. Accession numbers: *Sc* (NP_014489), *Pb*01 (PAAG_06164.1), *Pb*03 (PABG_06497.1), *Pb*18 (PADG_07957.1), *Cn* (CNAG_00876.2) and *Cg* (CNBG_6082.2).

siderophores. Acetyl is used for rhodotorulic acid and ferrichrome synthesis, while anhydromevalonyl is utilized in the fusarinines and coprogens pathway (Haas et al., 2008). A. fumigatus sidF encodes an N⁵-hydroxyornithine: cis anhydromevalonyl coenzyme A-N5-transacylase involved in the synthesis of fusarinine and triacetylfusarinine (Schrettl et al., 2007). The sidF ortholog of H. capsulatum, sid3 gene, is transcriptionally induced under iron restricted conditions (Hwang et al., 2008). Hydroxamates are covalently linked via peptide (rhodotorulic acid, ferrichromes, coprogens) or ester bonds (fusarinines, coprogens) carried out by non-ribosomal peptide synthetases (NRPSs; Finking and Marahiel, 2004). In A. fumigatus, sidC and sidD encode two NRPSs involved in ferricrocin (intracellular siderophore) and triacetylfusarinine C (TAFC) biosynthesis, respectively. Some siderophores additionally require acetylation at the N2-amino group, such as coprogen and TAFC. For example, sidG deletion in A. fumigatus results in the abrogation of the TAFC siderophore production (Schrettl et al., 2007). Given that our in silico analysis of P. brasiliensis identified sequences capable of coding for SidAp, SidFp, SidCp and SidDp, it is reasonable to hypothesize that P. brasiliensis may be able to synthesize both extracellular and intracellular siderophores.

Although *Cryptococcus* species have been described as unable to produce siderophores (Jacobson and Petro, 1987), *in silico* analysis of *C. neoformans* var. *grubii* and *C. gattii* structural genomes indicates the presence of *sidD* and *sidG* genes, which are also involved in other metabolic pathways in fungi. However, *sidA* and *sidF* genes were not found, and these genes are essential, especially since they act early in the pathway for siderophores production (**Table 2**; **Figure 1**). It will be interesting to examine if *sidA* and *sidF* have other functions and how siderophore-associated iron uptake was replaced to account for this loss.

Conserved domains in proteins related to siderophore biosynthesis

As described above, the third siderophore biosynthetic step is performed by NRPSs. These enzymes have a modular structure where one module, the catalytic unit, is composed of an adenylation domain (A) for substrate specificity and activation, a peptidyl carrier (PCP) domain that binds a 4'phosphopantetheine cofactor for attachment of the activated substrate, and a condensation (C) domain for bond formation (Finking and Marahiel, 2004). As *Cryptococcus* species are not siderophore producers, NRPSs domains analysis was performed only with SidCp ortholog found in *P. brasiliensis* genomes. These analyses revealed that, as in *A. fumigatus*, the three domains essential for NRPS function are present in SidCp from the three *P. brasiliensis* isolates examined (**Figure 3A**). Domains found in other siderophore biosynthesis related proteins are shown in **Table A2** in Appendix.

Siderophore uptake

The presence of orthologs for appropriate siderophore genes and the fact that the iron binding capacity of medium from low iron cultures of P. brasiliensis is greater than that of iron-replete medium (Arango and Restrepo, 1988) supports our hypothesis that P. brasiliensis produces and captures siderophores from the extracellular environment. Therefore, we have categorized putative P. brasiliensis siderophore transporters by sequence homology analysis (Table 2; Figure 1). Searches of the P. brasiliensis genomes revealed that all three isolates contain the S. cerevisiae gene homolog SIT sit1. S. cerevisiae can utilize siderophore-bound iron either by the reductive iron-assimilation system or by membrane transporters. In the latter case, the uptake is mediated by four transporters that differ in substrate specificity: Sit1p/Arn3p, Arn1p, Taf1p/Arn2p, Enb1p/Arn4p (Lesuisse et al., 1998; Heymann et al., 1999, 2000; Yun et al., 2000a,b). Sit1p/Arn3p recognizes ferrioxamines, coprogen, and ferrichromes lacking anhydromevalonic acid. Additionally, P. brasiliensis isolates possess the A. nidulans SIT gene homologs, mirB, and mirC (Table 2; Figure 1). Heterologs expression assays of A. nidulans mir genes in a S. cerevisiae mutant strain unable to uptake siderophores have demonstrated that MirBp transports native TAFC, a hydroxamate siderophore. The growth of P. brasiliensis is stimulated by coprogen B and dimerum acid (DA), a derivative of rhodotorulic acid from Blastomyces dermatitidis, suggesting that P. brasiliensis can use hydroxamate compounds as iron sources (Castaneda et al., 1988).

The siderophore transporter Sit1p/Arn3p and the transporters of the SIT-family (mirA, mirB and mirC) were found in C. neoformans var. grubii and C. gattii (Table 2; Figure 1). The homolog gene sit1/arn3 was previously identified in C. neoformans var. neoformans using SAGE employed to examine the transcriptome under ironlimiting and iron-replete conditions (Lian et al., 2005). Mutants defective in sit1 had increased melanin production and elevated transcript levels for the laccase gene, lac1. The melanin phenotype may be caused by changes in iron homeostasis or membrane trafficking, perhaps leading to altered copper loading of laccase in the cell wall. Studies with mutants lacking sit1/arn3 in C. neoformans var. grubii and C. neoformans var. neoformans have demonstrated that the gene sit1 is required for siderophore utilization (ferrioxamine B) and growth in low iron-environments (Tangen et al., 2007). An overview of the siderophore biosynthesis and uptake in *P. brasiliensis* and *Cryptococcus* species is shown in **Figure 1**.

Analysis of transmembrane domains in siderophore–iron transporters

Amino acid sequences of siderophore transporter orthologs found in *P. brasiliensis* isolates and *Cryptococcus* species were analyzed in the TopPred server to predict their transmembrane domain topologies. **Figure 3B** presents the transmembrane segments of Sit1p in *S. cerevisiae, P. brasiliensis* isolates, *C. neoformans* var. *grubii* and *C. gattii.*

Gene	Organism/ accession number	Predicted function	Orthologs in <i>Pb</i> 01, 03 and 18 (accession numbers)†	<i>E</i> -value*	Orthologs in <i>Cryptococcus</i> species (accession numbers) [†]	<i>E</i> -value*
sidA	A. fumigatus	Ornithine-N⁵-	PAAG_01682.1	0.0	Not identified	_
	XP_755103	monooxygenase	PABG_03730.1	0.0		
			PADG_00097.1	0.0		
sidF	A. fumigatus	N ⁵ -transacylases	PAAG_01680.1	0.0	Not identified	_
	XM_743567		PABG_03728.1	0.0		
			PADG_00100.1	0.0		
sidC	A. fumigatus	Non-ribosomal	PAAG_08527.1	0.0	Not identified	_
	XP_753088	peptide synthetase	PABG_04670.1	0.0		
			PADG_05295.1	0.0		
sidD	A. fumigatus	Non-ribosomal	PAAG_01679.1	0.0	CNAG_03588.2	e-40
	XP_748662	peptide synthetase	PABG_03726.1	0.0	CNBG_2041.2	e-41
			PADG_00102.1	0.0		
sidG	A. fumigatus	N ² -transacetylase	Not identified	-	CNAG_04355.2	2e-5
	XP_748685				CNBG_2703.2	e-6
sit1/arn3	S. cerevisiae	Siderophore	PAAG_06516.1	0.0	CNAG_00815.2	0.0
	NP_010849	transporter	PABG_02063.1	0.0	CNBG_1123.2	0.0
			PADG_00462.1	0.0		
mirA	A. nidulans	Siderophore	Not identified	-	CNAG_02083.2	0.0
	AY027565	transporter			CNBG_5232.2	0.0
mirB	A. nidulans	Siderophore	PAAG_01685.1	0.0	CNAG_07751.2	0.0
	XP_681809	transporter	PABG_03732.1	0.0	CNBG_2036.2	0.0
			PADG_00095.1	0.0		
mirC	A. nidulans	Siderophore	PAAG_02233.1	0.0	CNAG_07519.2	0.0
	AY135152	transporter	PABG_04747.1	0.0	CNBG_1087.2	e-44
			PADG_05373.1	0.0		

Table 2 | Orthologs to genes related to siderophore biosynthesis and to iron uptake by the non-reductive siderophore transport system in *P. brasiliensis* and *Cryptococcus* species.

*Similarities with E-values < 10⁻⁵ were considered significant.

*Accession numbers: PAAG refers to Pb01; PABG refers to Pb03; PADG refers to Pb18; CNAG refers to C. neoformans var. grubii and CNBG refers to C. gattii.

The number of segments varies between 12 and 15. Identical topology was found in Sit1p from *P. brasiliensis* isolates *Pb*03 and *Pb*18, whereas *Pb*01 has a different topology. Transmembrane domains were also identified in all the other siderophore transporters, as shown in **Table A2** in Appendix. These transporters also contain a MFS1 domain, which indicates that they belong to the MFS of transporters.

Iron source preferences

Several fungal pathogens utilize heme or hemoglobin as sources of iron (Foster, 2002; Jung et al., 2008). *C. albicans* expresses surface receptors for hemoglobin and hemolytic factors (Manns et al., 1994). Interestingly, heme–iron utilization in *C. albicans* is facilitated by Rbt5p, an extracellular glycosylphophatidylinositol (GPI)-anchored

hemoglobin-binding protein (Weissman et al., 2008). Although there is no experimental evidence regarding the utilization of iron from the heme group by *P. brasiliensis*, there are genes that show similarity with Hmx-1p (Pendrak et al., 2004), and exhibit a heme oxygenase domain (PAAG_06626.1 in *Pb*01; PABG_02644.1 in *Pb*03; PADG_01082.1 in *Pb*18) in each of the *P. brasiliensis* isolates. These genes are annotated as conserved hypothetical or as predicted proteins. *C. neoformans* var. *grubii* is also able to utilize heme and hemoglobin as iron sources, but the mechanism(s) of heme utilization by this fungus are still unclear (Jung et al., 2008).

Transferrin has also been shown to be an iron source for both *C. albicans* and *C. neoformans* var. *grubii*. These fungi employ high-affinity permeases to acquire iron from transferrin in mammalian



hosts through the reductive system (Knight et al., 2005; Jung et al., 2008). In the *P. brasiliensis* genome databases, genes were found (PAAG_04670.1; PABG_00038.1; PADG02428.1, respectively for isolates *Pb*01, *Pb*03 and *Pb*18) with high similarity to Cft1p, a permease from *C. neoformans* var. *grubii* required for iron utilization from transferrin (Jung et al., 2008).

COPPER

Copper uptake by the reductive system

Little is known about copper metabolism in P. brasiliensis. However, our in silico analyses of the S. cerevisiae copper metabolism-related genes in comparison to P. brasiliensis genomic databases revealed genes related to the copper reduction metalloreductase, fre. Copper transport is well described in S. cerevisiae where it is reduced from Cu (II) to Cu (I) by several cell surface metalloreductases encoded by several fre genes. These metalloreductases are regulated by iron and copper availability, mediated by the transcriptional factor Mac1p (Jungmann et al., 1993). Homologs of the copper metalloregulatory transcription factor gene (mac1) are present in both Pb01 and Pb03 genomes, but not in Pb18. Additionally, the high-affinity copper transport (Ctr3p) was found in all three isolate genomes. In S. cerevisiae, after reduction, copper is transported by the highaffinity copper transporter comprised by Ctr3p and Ctr1p, which are functionally redundant, although they have distinct amino acid sequences. Ctr3p is an integral membrane protein that assembles as a trimer to form a competent copper uptake permease at the plasma membrane. S. cerevisiae Ctr1p is localized at the plasma membrane and exists as an oligomer in vivo. These two high-affinity copper transport proteins are induced by copper deprivation and repressed by copper excess (Dancis et al., 1994a; Pena et al., 2000). In our in silico analyses, genes for the high-affinity copper transporter of the plasma membrane (ctr1) were not found, suggesting that highaffinity copper transport is performed only by the Ctr3p protein.

Genes related to metallochaperone (atx1), Cu²⁺ transporting P-type ATPase (ccc2) and superoxide dismutases (sod1 and sod2; Table 1) were also found in P. brasiliensis genomes. In the cell, copper is transported by Atx1p, a cytosolic copper metallochaperone protein, that transports Cu (I) to Ccc2p, a transporting P-type ATPase containing a cytoplasmic region containing two distinct soluble metal-binding domains that interact with Atx1p (Banci et al., 2007). Ccc2p mediates the export of copper from the cytosol and distributes it to cupric proteins (Yuan et al., 1997). S. cerevisiae also has a detoxification pathway formed by Cup1p and Cup2p, metallothioneins (Table 1), that protect against copper poisoning (Hamer et al., 1985). An alternative copper transport system is mediated by Ctr2p, a vacuolar membrane protein of S. cerevisiae, that mobilizes vacuolar copper stores to cytosolic copper chaperones (Rees et al., 2004). Homologs of the low-affinity copper transporter of the vacuolar membrane (Ctr2p) are in Pb03 and Pb18, but not in Pb01. Additionally, the metallothioneins (encoded by cup1 and cup2 genes) were not identified in *P. brasiliensis* isolates *Pb*01, *Pb*03 and *Pb*18.

In silico analysis (**Table 1**) revealed that *Cryptococcus* species have orthologs encoding ferric/cupric reductases, suggesting that the copper reduction process is similar to that described for *S. cerevisiae*. Homologs of the high-affinity copper transporter *ctr3* gene and copper metalloregulatory transcription factor gene (*mac1*) have previously been identified (Waterman et al., 2007). Also, proteins with similarity to the cytosolic copper metallochaperone (*atx1* gene), the Cu²⁺ transporting P-type ATPase (*ccc2* gene) and the cytosolic and mitochondrial superoxide dismutases (*sod1* and *sod2* genes) have also identified, suggesting that copper distribution in *Cryptococcus* species occurs as described in *S. cerevisiae*. A homolog of the *ctr2* gene was identified only in *C. neoformans* var. *grubii*. Recently it was demonstrated that Ctr2p links copper homeostasis to polysaccharide capsule production in *C. neoformans*. The lack of this protein resulted in increased phagocytosis by murine macro-

phage, sensitivity to copper starvation and defects in polysaccharide capsule formation and melanization (Chun and Madhani, 2010). The gene *ctr1* for the high-affinity copper transporter of the plasma membrane and the genes *cup1* and *cup2* for metallothioneins were not found in *Cryptococcus* species. These analyses suggest that the high-affinity copper transport in cryptococcal cells is primarily performed by the protein encoded by *ctr3*.

Analysis of conserved motifs present in copper transporters

Searches for conserved domains revealed the presence of Mets and MXXXM motifs in the Ctr3p of the P. brasiliensis isolates and the Cryptococcus species (Figure 4). Studies in yeast and mammalian cells have revealed that proteins of the CTR family are integral membrane proteins containing three membrane-spanning domains, with high protein sequence homology (Dancis et al., 1994a; Lee et al., 2002). With the exception of S. cerevisiae Ctr3p, all CTR family members are rich in methionine residues within the amino-terminal portion (Labbe et al., 1999). These residues are arranged as MXXM and/or MXM, called Mets motifs, and it has been suggested that they could be involved in extracellular copper binding (Dancis et al., 1994b). It has been demonstrated that these clustered methionine residues together with an MXXXM motif in the transmembrane domain of CTR family members are important for copper uptake (Puig et al., 2002). In P. brasiliensis the MXXXM motif is found within the third transmembrane segment. The Ctr3p of Cryptococcus species contains only two predicted transmembrane domains instead of the three transmembrane segments described for other fungi. In C. neoformans var. grubii and C. gattii, the MXXXM motif is within the second transmembrane domain. Conserved domains were also found in amino acid sequences of other proteins involved in copper metabolism (Table A1 in Appendix), suggesting that the orthologs found in P. brasiliensis and Cryptococcus may have activities that are similar to genes with established functions in other fungi.

ZINC

Zinc uptake

Comparisons to the *S. cerevisiae* genes related to zinc metabolism performed in *P. brasiliensis* genomes are presented in **Table 1**. Analyses demonstrate that *P. brasiliensis* has homologs to zinc trans-

porters described in S. cerevisiae that are localized in the plasmatic, vacuolar and endoplasmic reticulum membranes. Importantly, five genes encoding to transporters of the ZIP family, with homology to S. cerevisiae Zrt1p or Zrt2p, are in the P. brasiliensis genomic database. In S. cerevisiae, zinc is transported by proteins belonging to the ZIP family, which is composed by a zinc high-affinity transporter protein encoded by the zrt1 gene and a low-affinity transporter encoded by the zrt2 gene (Gaither and Eide, 2001). We have previously identified homologs of zinc transporters by transcriptional analysis of P. brasiliensis yeast cells after incubation in human blood and plasma (Bailão et al., 2006, 2007). Interestingly, P. brasiliensis isolate Pb01 has two vacuolar membrane zinc transporters, encoded by the zrc1 and cot1 genes, whereas isolates Pb03 and Pb18 contain only the cot1 homolog. Intracellularly, zinc is in vacuoles in association with the vacuolar membrane proteins Zrc1p and Cot1p, members of the cation diffusion facilitator (CDF) family (MacDiarmid et al., 2002). A homolog of the transcription factor Zap1p is also present in the three P. brasiliensis isolates. The expression of the genes associated with zinc homeostasis is positively regulated in S. cerevisiae by the transcription factor Zap1p, which regulates the expression of zrt1, zrt2, zrt3, fet4, and zcr1 under zinc limiting conditions (Wu et al., 2008). Therefore, zinc assimilation in *P. brasiliensis* may be similar to that of *S. cerevisiae*.

Similarly, zinc homeostasis in Cryptococcus species is poorly studied. In silico analysis was performed by comparing S. cerevisiae genes related to zinc metabolism in genomic cryptococcal databases (Table 1). The results show that C. neoformans var. grubii and C. gattii have Zrt1p and Zrt2p zinc transporters homologs. These proteins putatively internalize zinc into the cell. Further, homologs of the vacuolar transporter Cot1p and the CDF Msc2p are present. Cot1p is presumably in the vacuolar membrane and should be related to zinc storage in this compartment. Msc2p, an endoplasmic reticulum membrane zinc transporter, could be related to zinc transport to this organelle. The protein encoded by msc2 (CDF) is responsible for zinc homeostasis in the endoplasmic reticulum in S. cerevisiae (Ellis et al., 2004). A homolog of the transcription factor Zap1p is also present in *Cryptococcus*. Since homologs to the vacuolar membrane zinc transporter gene zrt3 were not identified, the zrc1 and cot1 genes, encoding vacuolar membrane zinc transporters



P. brasiliensis isolates and *Cryptococcus* species. Ctr3p from *P. brasiliensis* isolates contains three putative transmembrane domains (TMD1-3, shown in black) while Ctr3p from *Cryptococcus* species presents only two TMDs. All species contain putative copper binding motifs (Mets motifs) arranged as

MXXM and/or MXM. MXXXM motif in TMD3 in *P brasiliensis* isolates and TMD2 in *Cryptococcus* species are represented in white characters. The length of each protein, in amino acids, is shown on the right. Accession numbers: *Pb*01 (PAAG_05251.1), *Pb*03 (PABG_07607.1), *Pb*18 (PADG_05084.1), *Cn* (CNAG_00979.2) and *Cg* (CNBG_0560.2).

Pb03	MLGTYSHFTTLLLRRDEQENS	21
<i>Pb</i> 18	MGTYSHFTTLLLRRDEQENS	
Pb01	MLGTSSRLTTLLVRRDEQTNS	
Sc	MSNVTTPWWKQWDPSEVTLADKTPDDVWKTC	
Cn	MTTYRSILVYCVALTIVGLVHGHTTGEEEHDHDHDHQSEAVETDAAATDLHAG	
Cg	-MTVRSIFVYFVAFAVVGFVRGQIIAEEEHDHDHHDHHHDHHDHDHDDDAVVTDAATHDAHAG	
<i>Pb</i> 03		75
Pb18	PGACATGNEYDGHLNLRLSAVFVILIGSSIGALFPVWARPGRTNASKGRRVDVP	
Pb10 Pb01	PGACATGNEYDGHINLRLSAVFVILIGSSIGALFPVWARFGATNASAGAAVDVF	
Sc	VLQGVYFGGNEYNGNLGARISSVFVILFVSTFFTMFPLIST-KVKRLRIP	
Cn	HTHAHSSSDCWVTELONYDLSLHIAAVFVMVASAIGVFLPVIIGKLGSRNKLF	
Cg	HTHAHSSSDEWVIELQNIDLS	
<i>Pb</i> 03	PWAFFVAKYFGSGVIVATAFIHLLAPAHEALS-NPCLTGPVTEYPWVEGIMLMTIVLL	132
Pb18	PWAFFVAKYFGSGVIVATAFIHLLAPAHEALS-NPCLTGPVTEYPWVEGIMLMTIVLL	131
Pb01	PWA <mark>FFVAKYFGSGVIVATAFIHLLAP</mark> AHEALS-NPCLTGPV <mark>TEYPWVEGVMLMTIVLL</mark>	132
Sc	LYVYLFAKYFGSGVIVATAFIHLMDPAYGAIGGTTCVGQTGNW <mark>GLYSWCPAIMLTSLTFT</mark>	140
Cn	GS <mark>VFFVLKYFGSGIIISLAFVHLLI</mark> HAFFNLT-SECVGNLEYESA <mark>APAIAMATVIVV</mark>	163
Cg	GSLFSILKYFGSGIIISLAFVHLLIHAFFNLT-SACVGDMDYESVAPAIAMATVIVV	169
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Pb03	FFIELMAMRYARFGEADIAKELENGAWDMGHGHSHDNGHSNGKILAPNHTHTHTHDHDSV	192
Pb18	FFIELMAMRYARFGEADIAKELENGAWDMGHGHSHDHGHSNGKILAPNHTHTHDHDSV	191
<i>Pb</i> 01	FFIELMAMRYARFGEADIAKELENGAWDLCHCHSHDHCHSNGKILAPTHTHNHTNDHDSV	
SC	FLTDLESSVWVERKYGLSHDHTHDEIKDTVVRNTAAVSSENDNE	
Cn	WLVDFLGSRYITRQNSYVPECDRNISAALCSSSEPLGERK	
Cg	WLVDFFGSRYIARQNSCLPEGDRNITAASSSYPVSQGEKK	209
		and should be
Pb03	NSDVNTNIPGEDHLGHARHHLTDAVSKKNCHSFVGKTAADSKNHGPSDHIHGHMSLVEDY	252
Pb18	NSDVNTNFPGENHLGHARHHLTDAVSKKNCHSFVGKTAADSKNHGPPDHTHGHMSLVEDY	25.
Pb01	NSDVNIHMPGEDHLGHVRHHHTDAVSKKNCHSLVGKTAADSKNHSPPDRPHGHMALVEDY	
Sc	NGTANGSHDTKNGVEYYEDSDATSMDVVQSF	
Сп	KDDISTPMTELACCGPKNLEITNFDGAA	
Cg	IDGISTPMTELACCGPNKSKVTPFDGAA * ::	23
<i>Pb</i> 03	SAQLTSIFILEF <mark>C</mark> IIF <mark>HS</mark> VFI <mark>C</mark> LTLAVAGKEFITLYIVLVFHQTFEGLGLGARIATVPWP	312
<i>Pb</i> 18	SAQLT <mark>SIFILEFC</mark> IIF <mark>HS</mark> VFICLTLAVAGKEFITLYIVLVFHQTFEGLGLGARLATVPWP	311
Pb01	SAQLTSIFILEF <mark>GIIFHS</mark> VFI <mark>C</mark> LTLAVAGKEFITLYIVLVFHQTFEGLGLGARLATVPWP	312
Sc	QAQFYAFLILEF <mark>C</mark> VIF <mark>HS</mark> VMI <mark>G</mark> LNLGSVGDEFSSLYPVLVFHQSFEGLGIGARLSAIEFP	275
Cn	KTAHWNVQLLEY <mark>G</mark> VIE <mark>HS</mark> IMI <mark>G</mark> VSLGAMGTGFNTTFAALVFHQLFEGLGLGARIAMLIWP	293
Cg	KTAHWNVQLLEYCVIFHSIMICVSLGAMGTGFNTTFAALVFHQLFEGLGLGARIAMLVWP .: .**:*:****:**** * * * : .***** ********	29
<i>Pb</i> 03	GSKRLTPYILAIAFGLSTPVAIAIGLGVHETYPPESQTSLIVNGVFDSISAGILIYTA	370
Pb18	GSKRLTPYILAIAFGLSTPVAIAIGLGVHETYPPESQTSLIVNGVFDSISAGILIYTA	
Pb01	G-SKRLTPYILAIAFGLSTPIAIAIGLGIHETYPPESQTSLIVNGVFDSISAGILIYTA	
Sc	RSKRWWPWALCVAYGLTTPICVAIGLGVRTRYVSGSYTALVISGVLDAISAGILLYTG	
Cn	AGISSAIKKWSMCLAYALATPVGIAIGIGVHESVNMNGRAILLSTGILDSISAGILLYCG	
Cq	PGVSSTIKKWAMCLAYALVTPVGIAIGIGVHESINMNGRAILLSTGILDSISAGILLYGG	
~9	*. :* ** : **** : *** · * · · * · · * · · * · · * · · * · · * · · * · · * · · * · · * ·	55
Pb03	LVELMAHEFMFSTTMRRASIRTVLAAFALLCLGAALMALLGKWA 414	
Pb18	LVELMAHEFMFSTTMRRASIRTVLAAFALLCLGAALMALLGKWA 413	
Pb01	LVELMAHEFMFSTTMRRASIRTVLAAFALLCLGAALMALLGKWA 414	
Sc	LVELLARDFIFNP-QRTKDLRELSFNVICTLFGAGIMALIGKWA 376	
Cn	LCQLLYREWVVGE-MRDASTSKIIVALVSLFLGLFAMSFIGKWI 394	
Cg	LCOLLYREWVVGE MRDASTGEIIVALVSLFLGLFAMSFIGKWI 400	
-9	* :*: :::: * . : . :* *:::***	

FIGURE 5 | Ali P. brasiliensis isolates and Cryptococcus species. The predicted

transmembrane domains are shown in gray boxes. The black boxes inside the transmembrane segment contain conserved histidine-serine and glycine residues. The histidines found in the amino-terminal region of Zrt1p from

nains III and IV in P. brasiliensis and S. cerevisiae are boxed. Asterisks indicate amino acid identity and dots represent conserved substitutions. Accession numbers: Pb03 (PABG_07725.1), Pb18 (PADG_08567.1), Pb01 (PAAG_08727.1), Sc (NP_011259), Cn (CNAG_03398.2) and Cg (CNBG_2209.2).

could be responsible for the zinc transport to this organelle. This analysis suggests that *C. neoformans* var. *grubii* and *C. gattii* could obtain zinc via routes similar to that described for *S. cerevisiae*.

Analysis of conserved regions in the high-affinity zinc transporter (Zrt1p) in P. brasiliensis isolates and Cryptococcus species

Alignment of Zrt1p amino acid sequence from S. cerevisiae, P. brasiliensis isolates and Cryptococcus species revealed some conserved features (Figure 5). Concerning the predicted transmembrane domain number, all P. brasiliensis isolates contain eight predicted domains, while both C. neoformans var. grubii and C. gattii have nine. Proteins belonging to the ZIP family are predicted to have from five to eight transmembrane domains and they vary in size from 233 to 477 amino acid residues. The variations in the aminoterminal portion are usually responsible for the differences in size. The transmembrane domain IV has the most conserved portions of ZIP family proteins, with conserved histidine and glycine residues. The histidine residue and the adjacent polar residue, usually a serine, within the transmembrane domain are predicted to comprise part of a heavy metal-binding site in the center of the membrane (Eng et al., 1998). The amino acid sequence of S. cerevisiae Zrt1p presents a number of histidine residues in a large loop between the transmembrane segments III and IV, which is a putative metal ion binding site (Zhao and Eide, 1996a). The histidine-serine and glycine residues are conserved within the fourth transmembrane region in P. brasiliensis and within the fifth transmembrane region in Cryptococcus. Regarding the histi-

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dine rich region, it is conserved between transmembrane domains III and IV in *P. brasiliensis* isolates, whereas are conserved at the amino-terminal portion in *Cryptococcus* species, as occurs in other members of the ZIP family (Eng et al., 1998). Conserved domains are also found in amino acid sequences of other proteins involved in zinc metabolism that were identified in the search for orthologs (**Table A1** in Appendix).

CONCLUSION

As we have described, microorganisms are extremely well equipped to exploit host metal sources during growth and infection. *Cryptococcus* species demonstrate remarkable flexibility in gaining access to and utilizing iron, the most investigated micronutrient in this organism. Our laboratories have begun to elucidate the mechanisms for the uptake and metabolism of micronutrients such as iron, copper and zinc in *P. brasiliensis*. Studies on individual genes and pathways are revealing unique features of micronutrients metabolism in this fungus. The application of systems biology approaches that incorporates genomic and proteomic data will further generate hypotheses about the common and specific responses to micronutrient deprivation in both pathogenic fungi and potentially lead to the development of novel therapeutics exploiting their metal requirements.

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APPENDIX

Table A1 | Conserved domains in proteins involved in iron, copper and zinc uptake by reductive systems in *P. brasiliensis* isolates and *Cryptococcus* species.

Gene product	Predicted function	Organism/accession number [†]	Conserved domains*	Transmembrane domains*	Signal peptide*
Fre1	Metalloreductase	P. brasiliensis 01/PAAG_05370.1	Ferric reductase domain	7	Yes
		P. brasiliensis 03/PABG_06003.1	FAD-binding domain	6	No
			NAD-binding domain		
Fre3	Metalloreductase	P. brasiliensis 01/PAAG_02079.1	Ferric reductase domain	6	Yes
		P. brasiliensis 03/PABG_02329.1	FAD-binding domain	6	Yes
		P. brasiliensis 18/PADG_00813.1	NAD-binding domain	6	Yes
Fre5	Metalloreductase	P. brasiliensis 03/PABG_07812.1	Ferric reductase domain	6	No
			FAD-binding domain		
			NAD-binding domain		
Fre7	Metalloreductase	P. brasiliensis 01/PAAG_06164.1		8	No
		P. brasiliensis 03/PABG_06497.1	Ferric reductase domain	8	No
		P. brasiliensis 18/PADG_07957.1	FAD-binding domain	8	No
		C. neoformans/CNAG_00876.2	NAD-binding domain	7	No
		C. gattii/CNBG_6082.2		8	No
Fre8	Metalloreductase	C. neoformans/CNAG_07334.2	Ferric reductase domain	6	No
			FAD-binding domain		
		C. gattii/CNBG_2116.2	NAD-binding domain	6	No
Fre10	Metalloreductase	C. neoformans/CNAG_06821.2	Ferric reductase domain	4	No
			FAD-binding domain		
		C. gattii/CNBG_5888.2	NAD-binding domain	4	No
Cfl4	Metalloreductase	C. neoformans/CNAG_06524.2	Ferric reductase domain	5	No
			FAD-binding domain		
			NAD-binding domain		
Frp1	Metalloreductase	P. brasiliensis 01/PAAG_04493.1	Ferric reductase domain	5	No
		P. brasiliensis 03/PABG_04278.1	FAD-binding domain	6	No
		P. brasiliensis 18/PADG_04652.1	NAD-binding domain	5	No
Fet3	Ferroxidase	C. neoformans CNAG_06241.2	Copper-oxidase domain	1	Yes
Fet5	Ferroxidase	P. brasiliensis 03/PABG_05667.1	Copper-oxidase domain	_	No
		P. brasiliensis 18/PADG_05994.1		_	No
		C. neoformans/CNAG_07865.2		1	Yes
		C. gattii/CNBG_4942.2		1	Yes
Fet31	Ferroxidase	P. brasiliensis 01/PAAG_06004.1	Copper-oxidase domain	1	No
		C. neoformans/CNAG_02958.2		_	Yes
Fet33	Ferroxidase	P. brasiliensis 01/PAAG_00163.1	Copper-oxidase domain	_	No
		P. brasiliensis 03/PABG_05183.1		_	Yes
Ftr1/Ftr2	Iron permease	C. neoformans/CNAG_06242.2	FTR1 domain	7	Yes
		C. gattii/CNBG_3602.2		6	Yes
Fth1	Iron permease	C. neoformans/CNAG_02959.2	FTR1 domain	7	Yes
		<i>C. gattii/</i> CNBG_4943.2		7	Yes

Table A1 | Continued

Gene product	Predicted function	Organism/accession number [†]	Conserved domains*	Transmembrane domains*	Signal peptide*
Smf1	Low-affinity Permease	C. neoformans/CNAG_05640.2	Nramp domain	11	No
		C. gattii/CNBG_6162.2		11	No
Ccc1	Vacuolar transporter	P. brasiliensis 01/PAAG_07762.1	DUF125 domain	4	No
		P. brasiliensis 03/PABG_00362.1		4	No
		P. brasiliensis 18/PADG_02775.1		4	No
		C. neoformans/CNAG_05154.2		4	No
		C. gattii/CNBG_4540.2		4	No
Mrs3/Mrs4	Mitochondrial iron	P. brasiliensis 01/PAAG_05053.1	Mitochodrial carrier	_	No
	transporter	P. brasiliensis 03/PABG_04509.1	domain	-	No
		P. brasiliensis 18/PADG_04903.1		-	No
		C. neoformans/CNAG_02522.2		-	No
		C. gattii/CNBG_4218.2		-	No
Yfh1	Mitochondrial matrix iron	P. brasiliensis 01/PAAG_02608.1	Frataxin domain	-	No
	chaperone	P. brasiliensis 03/PABG_03095.1		-	No
		P. brasiliensis 18/PADG_01626.1		-	No
		C. neoformans/CNAG_05011.2		-	No
		C. gattii/CNBG_4670.2		_	No
Ggt1	Secreted glutathione-	P. brasiliensis 01/PAAG_06130.1	Gamma-	1	Yes
	dependent ferric reductase	P. brasiliensis 03/PABG_06527.1	glutamyltranspeptidase	1	Yes
		P. brasiliensis 18/PADG_07986.1	domain	1	Yes
		C. neoformans/CNAG_02888.2		_	No
		C. gattii/CNBG_3537.2		-	No
Mac1	Copper metalloregulatory	P. brasiliensis 01/PAAG_08210.1	Copper fist domain	-	No
	transcription factor	P. brasiliensis 03/PABG_07429.1		-	No
		C. neoformans/CNAG_07724.2		_	No
		C. gattii/CNBG_2252.2		_	No
Ctr3	High-affinity copper	P. brasiliensis 01/PAAG_05251.1	Ctr domain	3	No
	transporter of the plasma	P. brasiliensis 03/PABG_07607.1		3	No
	membrane	P. brasiliensis 18/PADG_05084.1		3	No
		C. neoformans/CNAG_00979.2		2	No
		C. gattii/CNBG_0560.2		2	No
Ctr2	Putative low-affinity copper	P. brasiliensis 03/PABG_01536.1	Ctr domain	3	No
	transporter of the vacuolar	P. brasiliensis 18/PADG_04146.1		3	No
	membrane	C. neoformans/CNAG_01872.2		3	No
Atx1	Cytosolic copper	P. brasiliensis 01/PAAG_00326.1	HMA domain	_	No
	metallochaperone	P. brasiliensis 03/PABG_06615.1		_	No
		P. brasiliensis 18/PADG_02352.1		_	No
		C. neoformans/CNAG_02434.2		_	No
		<i>C. gattii/</i> CNBG_4136.2		_	No
Ccc2	Cu ²⁺ transporting P-type	P. brasiliensis 01/PAAG_07053.1		7	No
	ATPase	P. brasiliensis 03/PABG_03057.1	HMA domain	8	No

Table A1 | Continued

Gene product	Predicted function	Organism/accession number [†]	Conserved domains*	Transmembrane domains*	Signal peptide*
		P. brasiliensis 18/PADG_01582.1	Hydrolase domain	8	No
		C. neoformans/CNAG_06415.2	E1-E2 ATPase domain	8	No
		C. gattii/CNBG_5045.2		8	No
Sod1	Cytosolic superoxide	P. brasiliensis 01/PAAG_04164.1	SOD domain	-	No
	dismutase	P. brasiliensis 03/PABG_03954.1		-	No
		P. brasiliensis 18/PADG_07418.1		-	No
		C. neoformans/CNAG_01019.2		-	No
		C. gattii/CNBG_0599.2		_	No
Sod2	Mitochondrial superoxide	P. brasiliensis 01/PAAG_02725.1	SOD N-terminal domain	-	No
	dismutase	P. brasiliensis 03/PABG_03204.1	SOD C-terminal domain	-	No
		P. brasiliensis 18/PADG_01755.1		-	No
		C. neoformans/CNAG_04388.2		_	No
		C. gattii/CNBG_2661.2		_	No
Zrt1	High-affinity zinc	P. brasiliensis 01/PAAG_08727.1	Zip domain	8	No
	transporter of the plasma	P. brasiliensis 03/PABG_07725.1		8	No
	membrane	P. brasiliensis 18/PADG_08567.1		8	No
		C. neoformans/CNAG_03398.2		9	Yes
		C. gattii/CNBG_2209.2		9	Yes
Zrt2	Low-affinity zinc	P. brasiliensis 01/PAAG_03419.1	Zip domain	8	Yes
	transporter of the plasma	P. brasiliensis 03/PABG_05498.1		7	No
	membrane	P. brasiliensis 18/PADG_06417.1		8	Yes
		C. neoformans/CNAG_00895.2		8	Yes
Zrc1	Vacuolar membrane zinc transporter	P. brasiliensis 01/PAAG_00702.1	Cation efflux domain	6	Yes
Cot1	Vacuolar membrane zinc transporter	P. brasiliensis 01/PAAG_07885.1	Cation efflux domain	5	Yes
		P. brasiliensis 03/PABG_07467.1		4	No
		P. brasiliensis 18/PADG_08196.1		5	Yes
		C. neoformans/CNAG_02806.2		6	Yes
		C. gattii/CNBG_3460.2		4	Yes
Zrt3	Vacuolar membrane zinc	P. brasiliensis 01/PAAG_09074.1	Zip domain	6	No
	transporter	P. brasiliensis 03/PABG_04697.1		6	No
		P. brasiliensis 18/PADG_05322.1		6	No
Msc2	Cation diffusion facilitator	P. brasiliensis 03/PABG_07115.1	Cation efflux domain	10	No
	protein of the endoplasmic	P. brasiliensis 18/PADG_06381.1		10	No
	reticulum and nucleus	C. neoformans/CNAG_05394.2		11	No
		C. gattii/CNBG_4458.2		10	No
Zap1	Zinc-regulated	P. brasiliensis 01/PAAG_03645.1	Zinc finger C ₂ H ₂ domain	-	No
	transcription factor	P. brasiliensis 03/PABG_03305.1	~ ~	_	No
		P. brasiliensis 18/PADG_01870.1		-	No
		C. neoformans/CNAG_05392.2		-	No
		C. gattii/CNBG_4460.2		_	No

*Amino acid sequence analysis was performed using the online software SMART.

[†]Accession numbers: PAAG refers to Pb01; PABG refers to Pb03; PADG refers to Pb18; CNAG refers to C. neoformans var. grubii and CNBG refers to C. gattii.

Gene product	Predicted function	Organism/accession number [†]	Conserved domains*	Transmembrane domains*	Signal peptide*
SidA	Ornithine-N ⁵ -	P. brasiliensis 01/PAAG_01682.1	Pyr_redox_2 domain	_	No
	monooxygenase	P. brasiliensis 03/PABG_03730.1		_	No
		P. brasiliensis 18/PADG_00097.1		_	No
SidF	N⁵-transacylases	P. brasiliensis 01/PAAG_01680.1	AlcB domain	_	No
		P. brasiliensis 03/PABG_03728.1		_	No
		P. brasiliensis 18/PADG_00100.1		_	No
SidC	Non-ribosomal	P. brasiliensis 01/PAAG_08527.1	Adenylation domain	_	No
	peptide synthetase	P. brasiliensis 03/PABG_04670.1	Peptidyl carrier domain	_	No
		P. brasiliensis 18/PADG_05295.1	Condensation domain	_	No
SidD	Non-ribosomal	P. brasiliensis 01/PAAG_01679.1	Adenylation domain	_	Yes
	peptide synthetase	P. brasiliensis 03/PABG_03726.1	Peptidyl carrier domain	_	No
		P. brasiliensis 18/PADG_00102.1		_	No
		C. neoformans/CNAG_03588.2	Condensation domain	_	No
		C. gattii/CNBG_2041.2		_	No
SidG	N²-transacetylase	C. neoformans/CNAG_04355.2	MYND-type zinc finger domains	-	No
		C. gattii/CNBG_2703.2	Acetyltransferase domain	_	No
Sit1/Arn3	Siderophore	P. brasiliensis 01/PAAG_06516.1	MFS1 domain	12	No
	transporter	P. brasiliensis 03/PABG_02063.1		14	No
		P. brasiliensis 18/PADG_00462.1		14	No
		C. neoformans/CNAG_00815.2		13	No
		C. gattii/CNBG_1123.2		13	No
MirA	Siderophore	C. neoformans/CNAG_02083.2	MFS1 domain	12	No
	transporter	C. gattii/CNBG_5232.2		11	No
MirB	Siderophore	P. brasiliensis 01/PAAG_01685.1	MFS1 domain	14	No
	transporter	P. brasiliensis 03/PABG_03732.1		14	No
		P. brasiliensis 18/PADG_00095.1		14	No
		C. neoformans/CNAG_07751.2		14	No
		C. gattii/CNBG_2036.2		14	No
MirC	Siderophore	P. brasiliensis 01/PAAG_02233.1	MFS1 domain	8	No
	transporter	P. brasiliensis 03/PABG_04747.1		12	No
		P. brasiliensis 18/PADG_05373.1		12	No
		C. neoformans/CNAG_07519.2		10	No
		C. gattii/CNBG_1087.2		14	Yes

Table A2 | Conserved domains in proteins related to siderophore biosynthesis and to iron uptake by the non-reductive siderophore transport system in *P. brasiliensis* isolates and *Cryptococcus* species.

*Amino acid sequence analysis was performed using the online software SMART.

[†]Accession numbers: PAAG refers to Pb01; PABG refers to Pb03; PADG refers to Pb18; CNAG refers to C. neoformans var. grubii and CNBG refers to C. gattii.